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(54) **TISSUE RESTORATION DEVICES, SYSTEMS, AND METHODS**

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CPC **A61F 5/0013** (2013.01); **A61F 5/0086** (2013.01); **A61B 17/122** (2013.01); **A61B 17/1227** (2013.01)

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CPC A61B 17/122; A61B 17/1227; A61B 17/1285; A61B 17/122804; A61B 17/2816; A61B 17/08; A61B 17/083; A61B 17/0487; A61B 2017/12004; A61B 2017/081; A61F 2005/414; A61F 5/005; A61F 2/2481; Y10S 128/25; A45D 8/24; A45D 8/20
USPC 606/139, 142, 144, 148, 151, 157–158, 606/120; 24/326, 335–341, 67.3, 67.5, 558, 24/565, 567; 251/4, 7, 9, 10, 117
See application file for complete search history.

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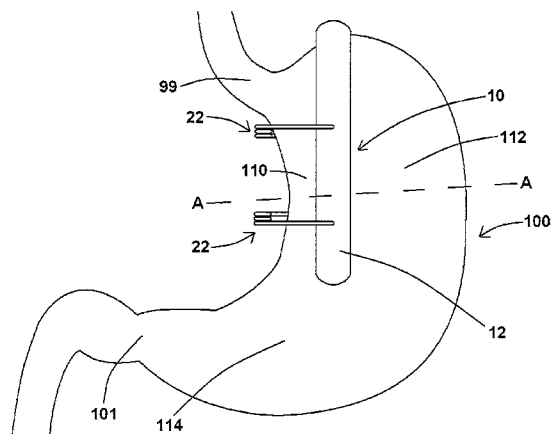
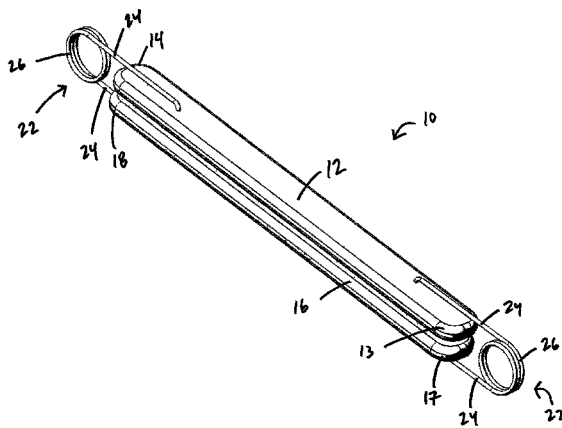
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(57) **ABSTRACT**

Tissue restoration devices, systems, and methods. In at least one embodiment, an implantable restraining device of the present disclosure comprises a first engaging component and a second engaging component, each component configured for laparoscopic insertion into a body cavity, a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, and a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend.

32 Claims, 34 Drawing Sheets



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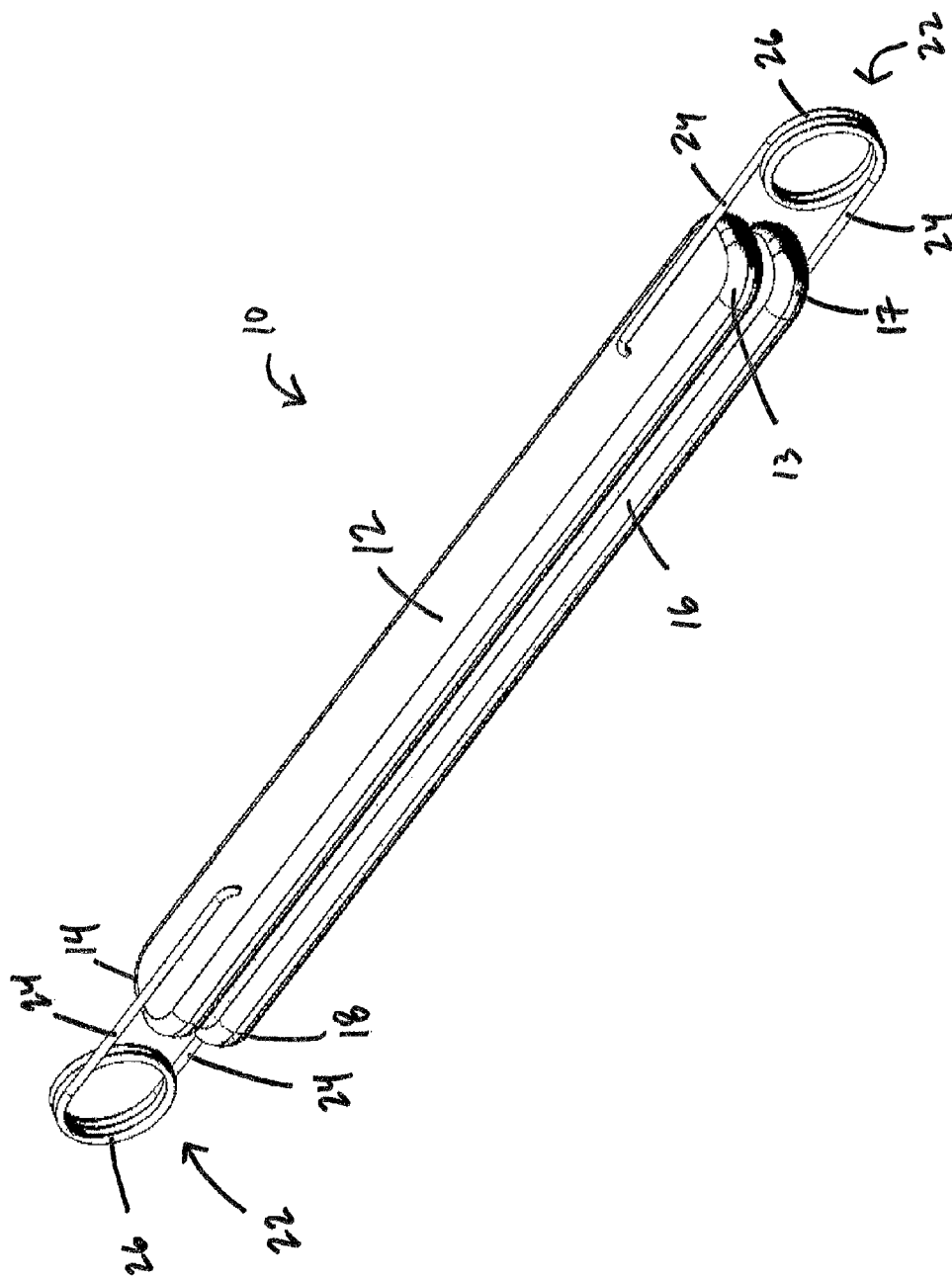


Fig. 1

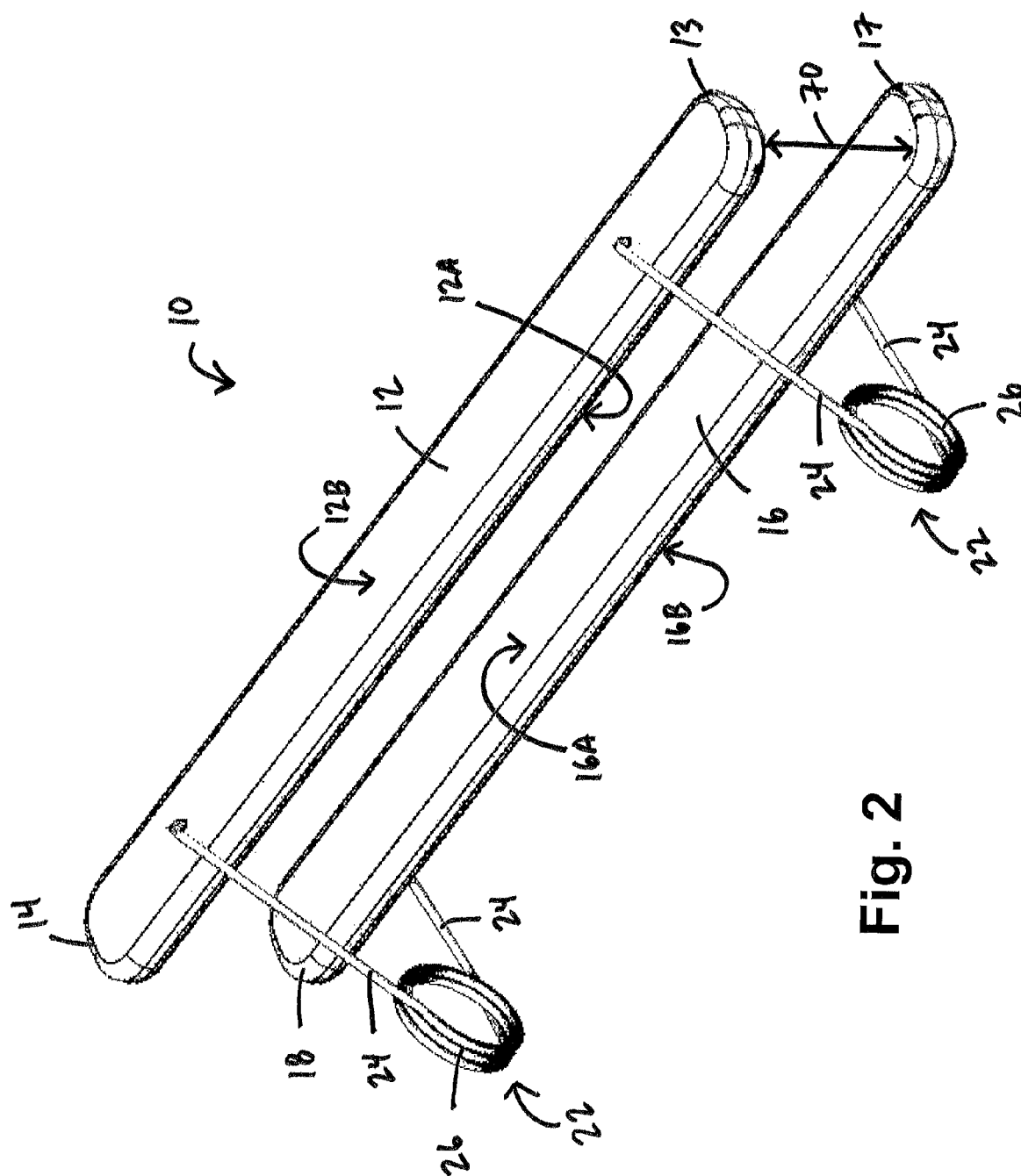


Fig. 2

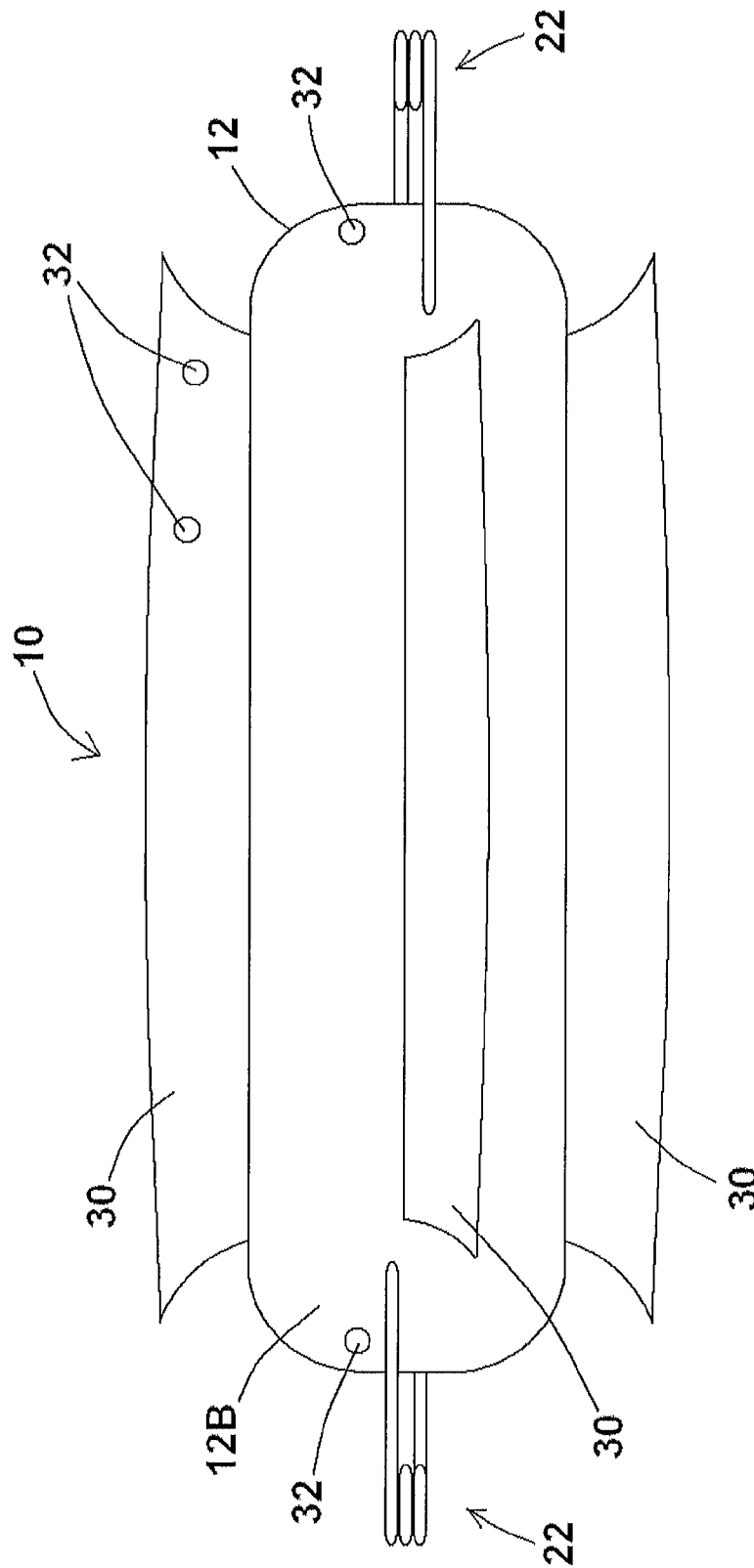
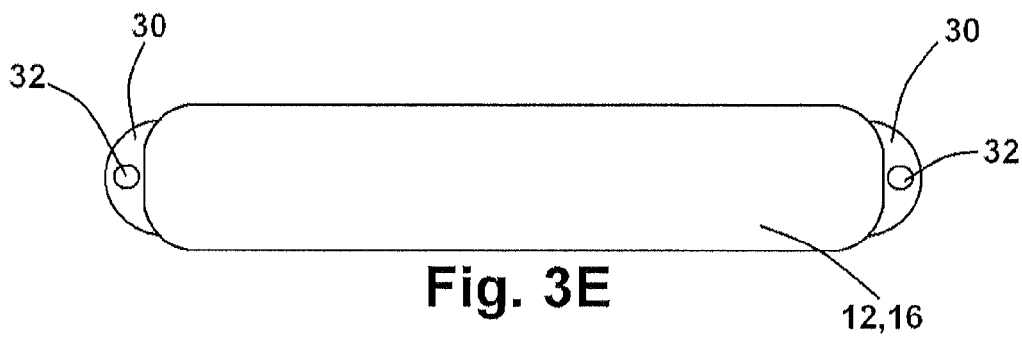
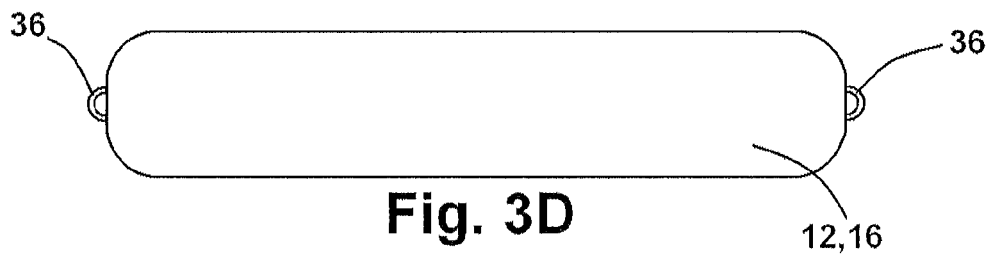
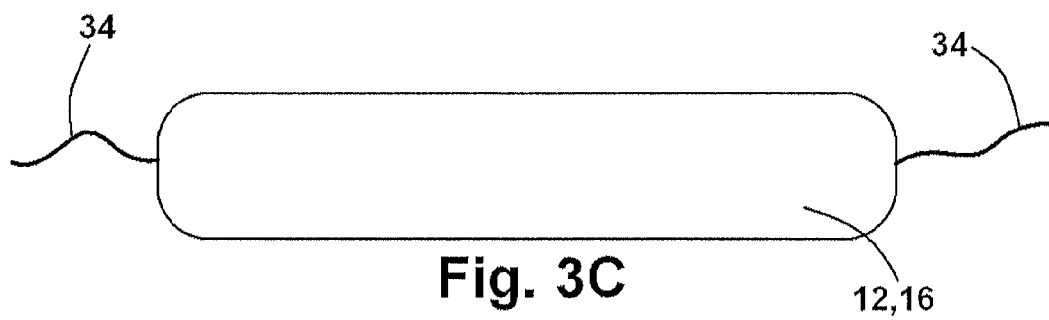
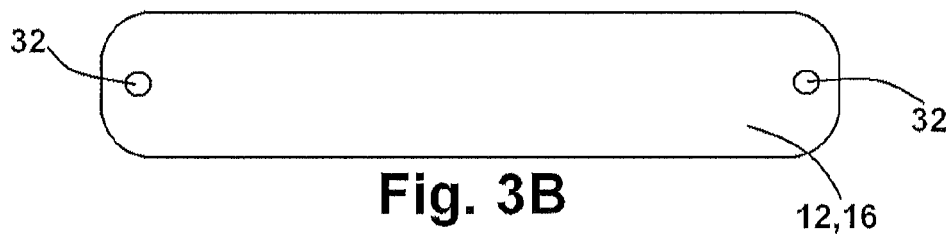


Fig. 3A



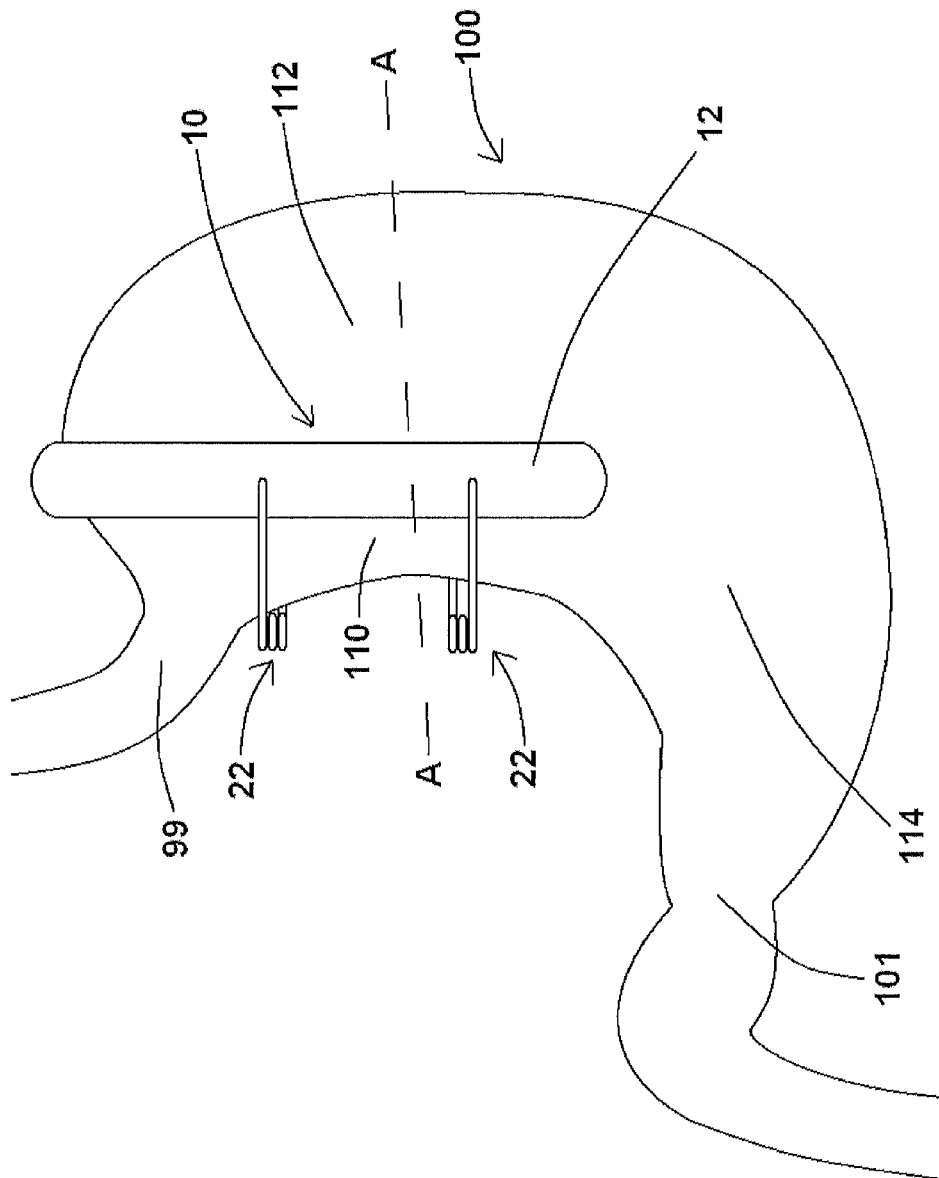


Fig. 4A

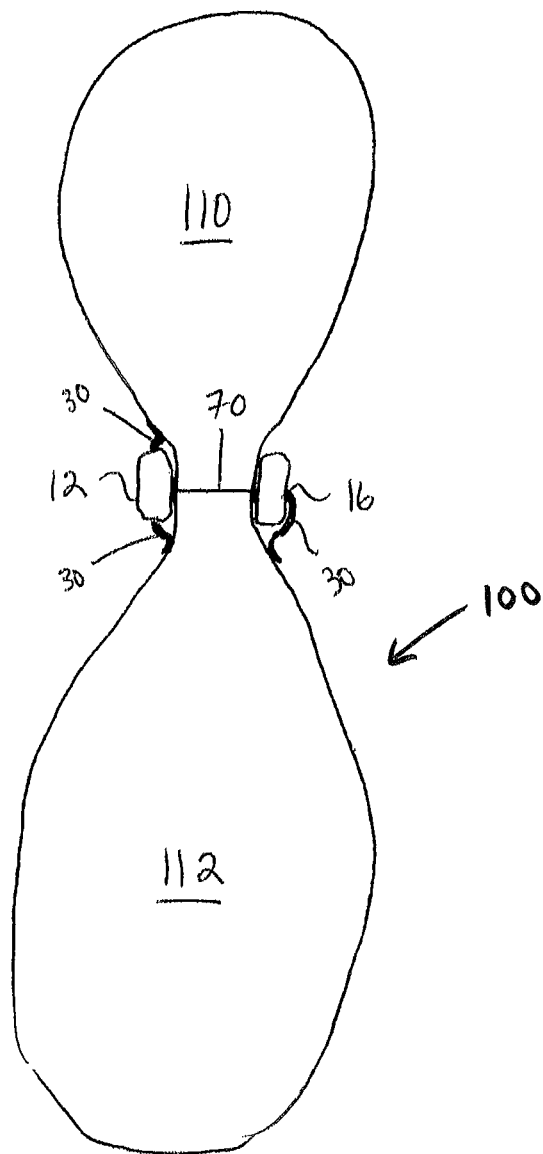


Fig. 4B

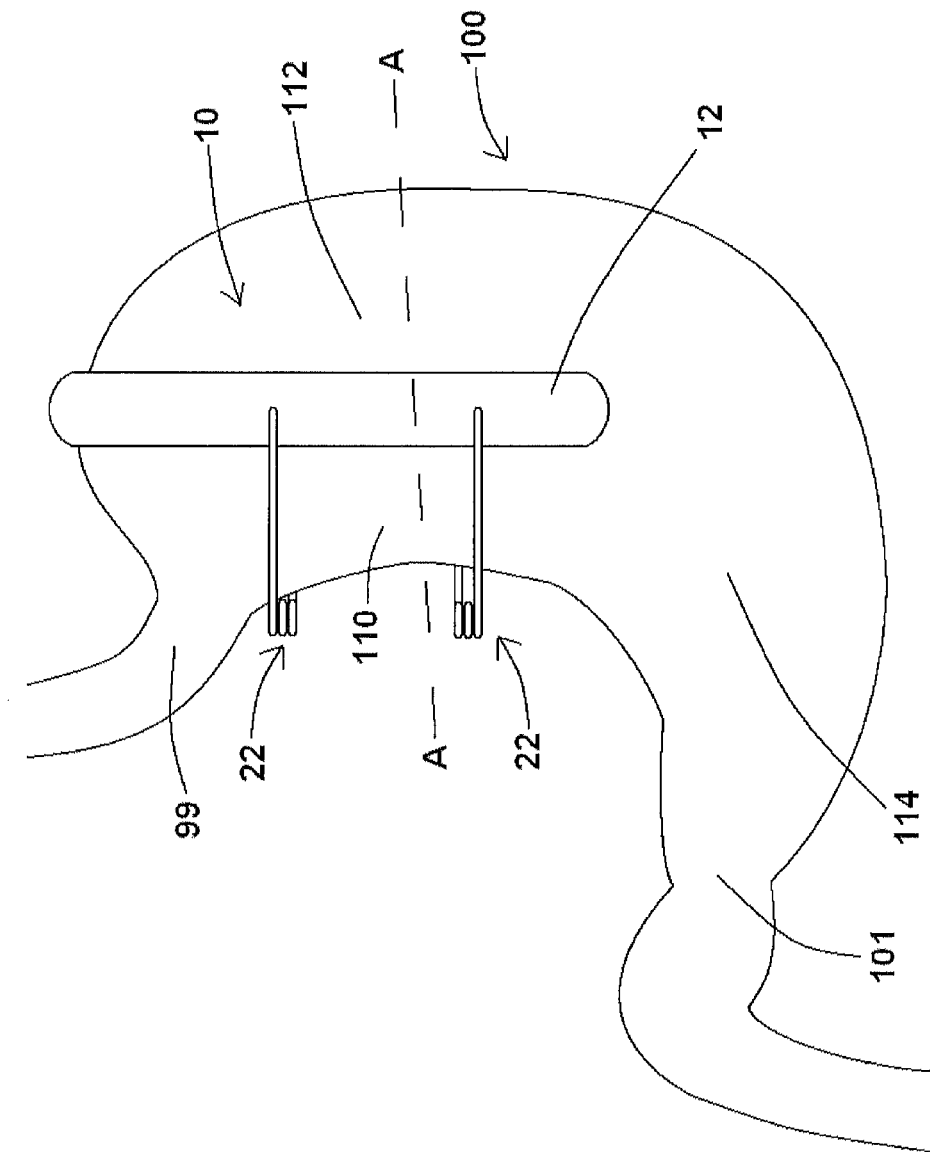


Fig. 5A

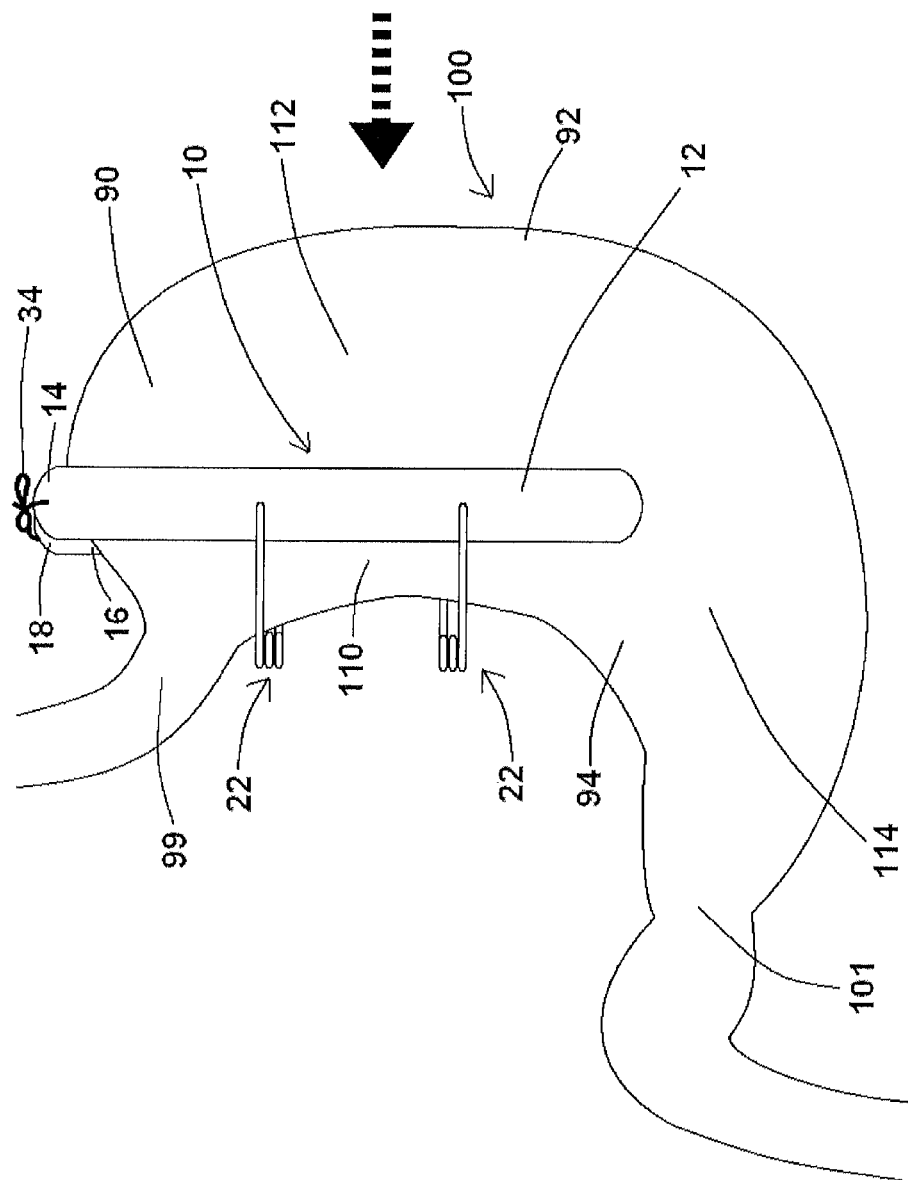


Fig. 5B

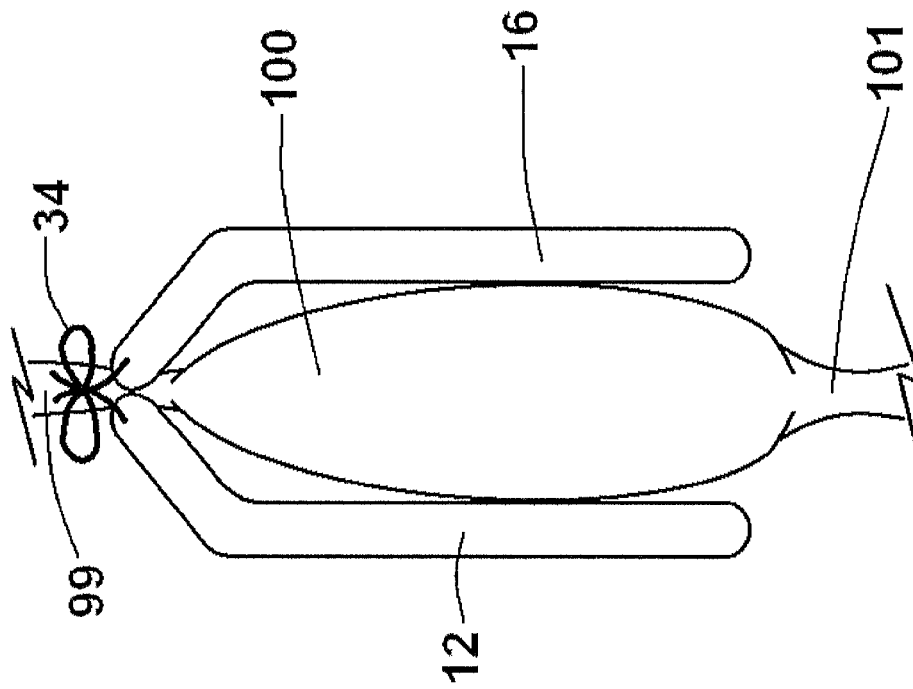


Fig. 5C

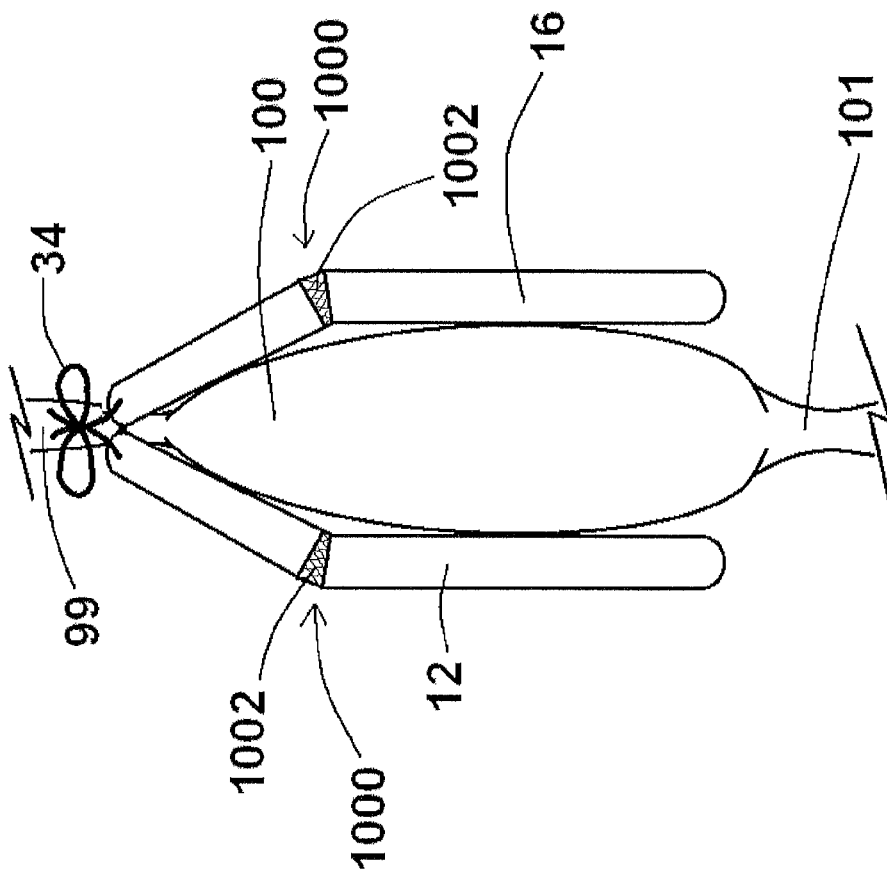
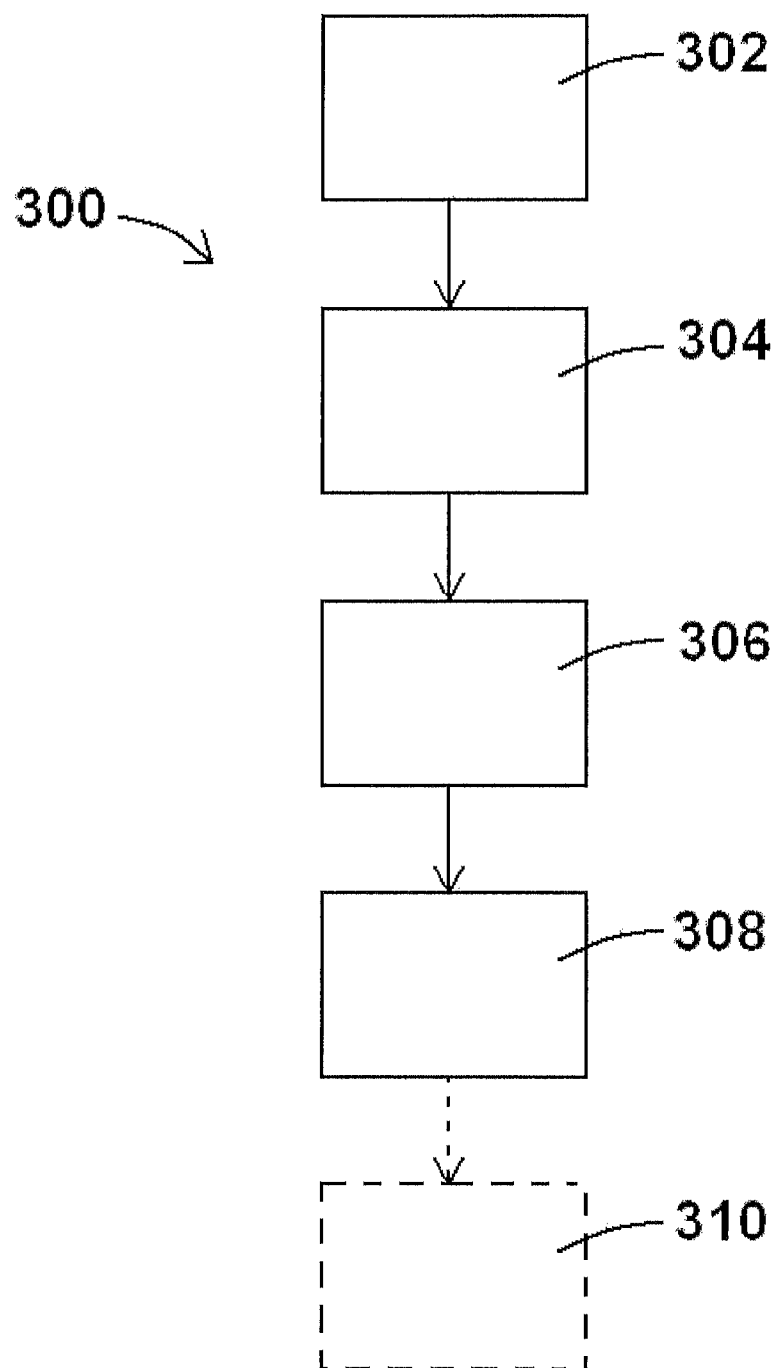
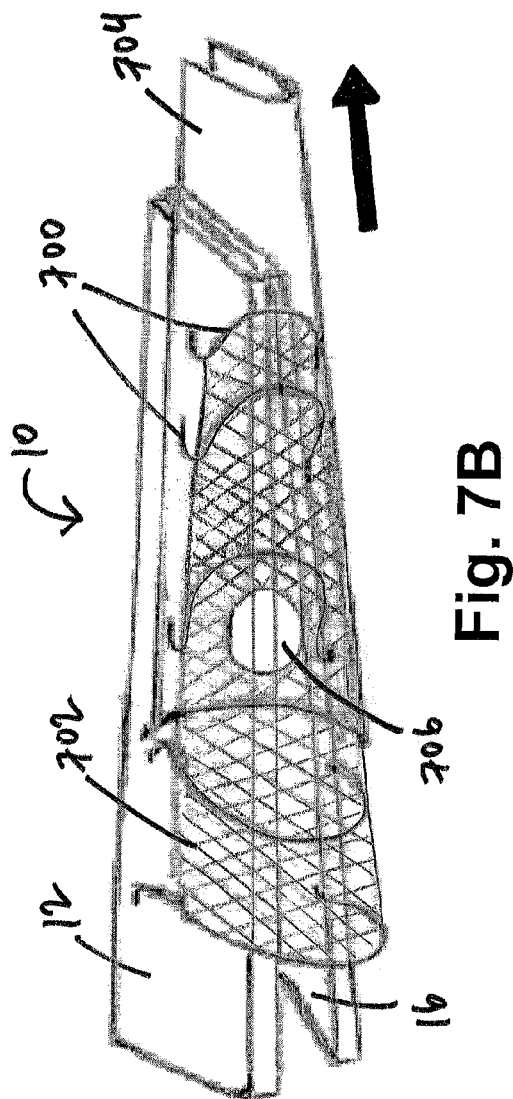
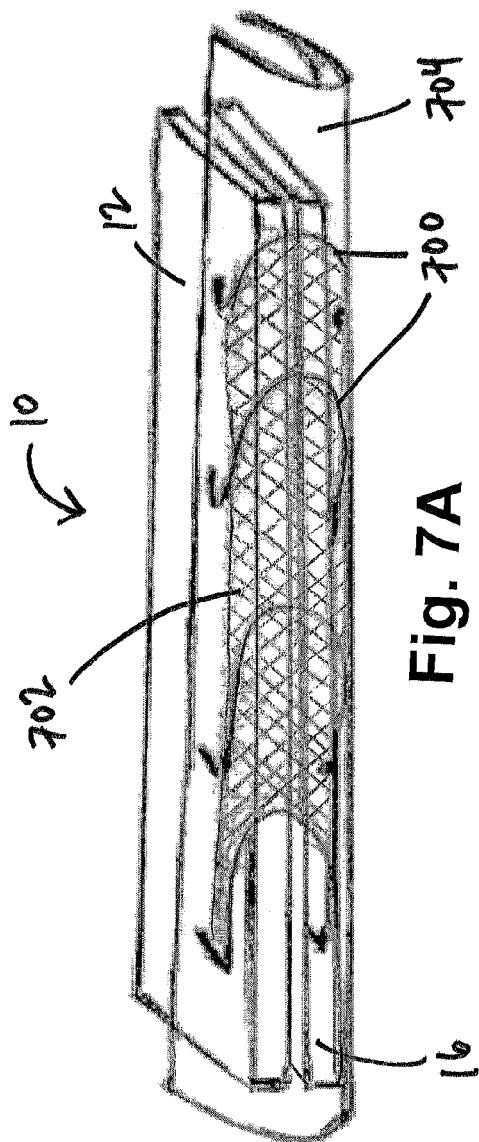


Fig. 5D

**Fig. 6**



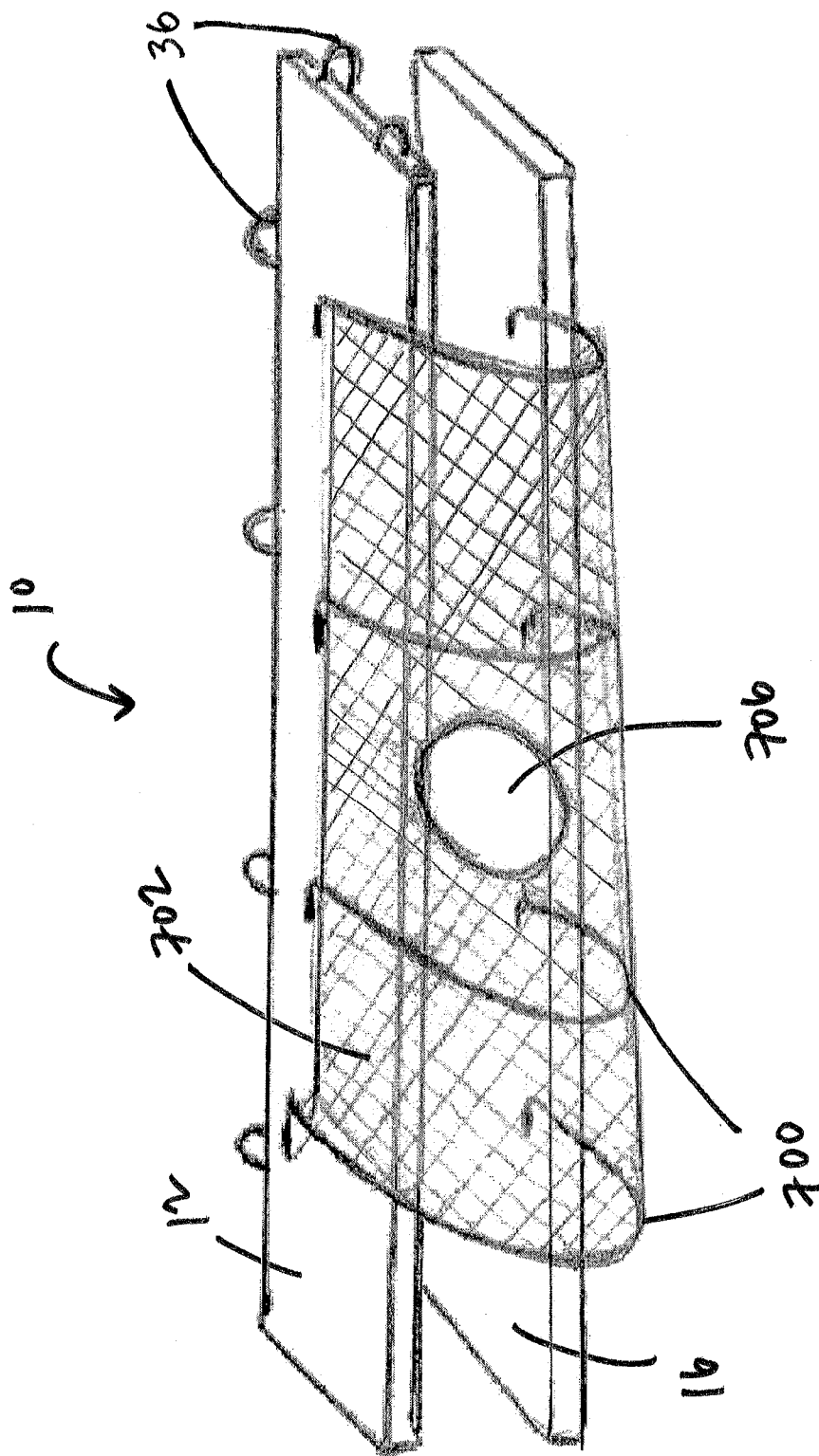


Fig. 8

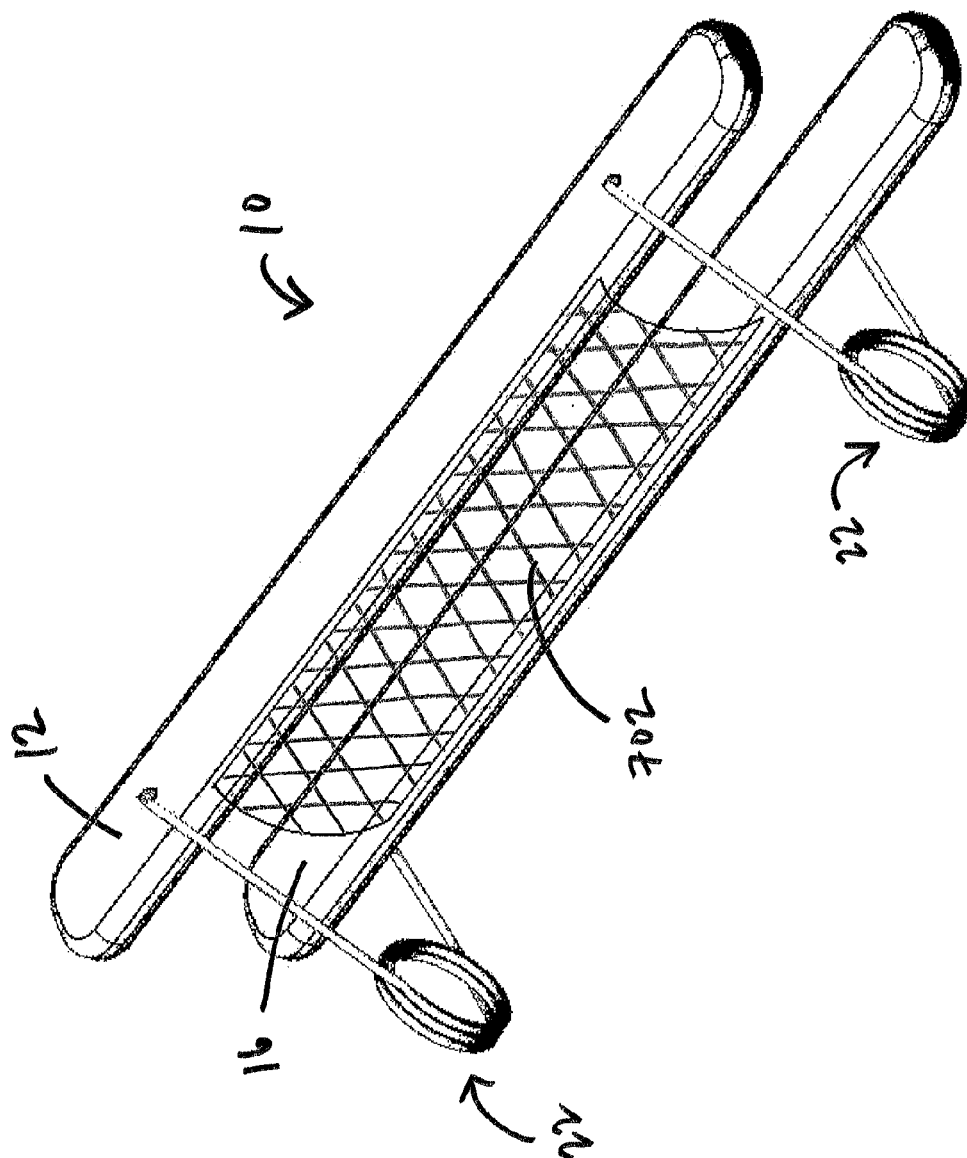


Fig. 9

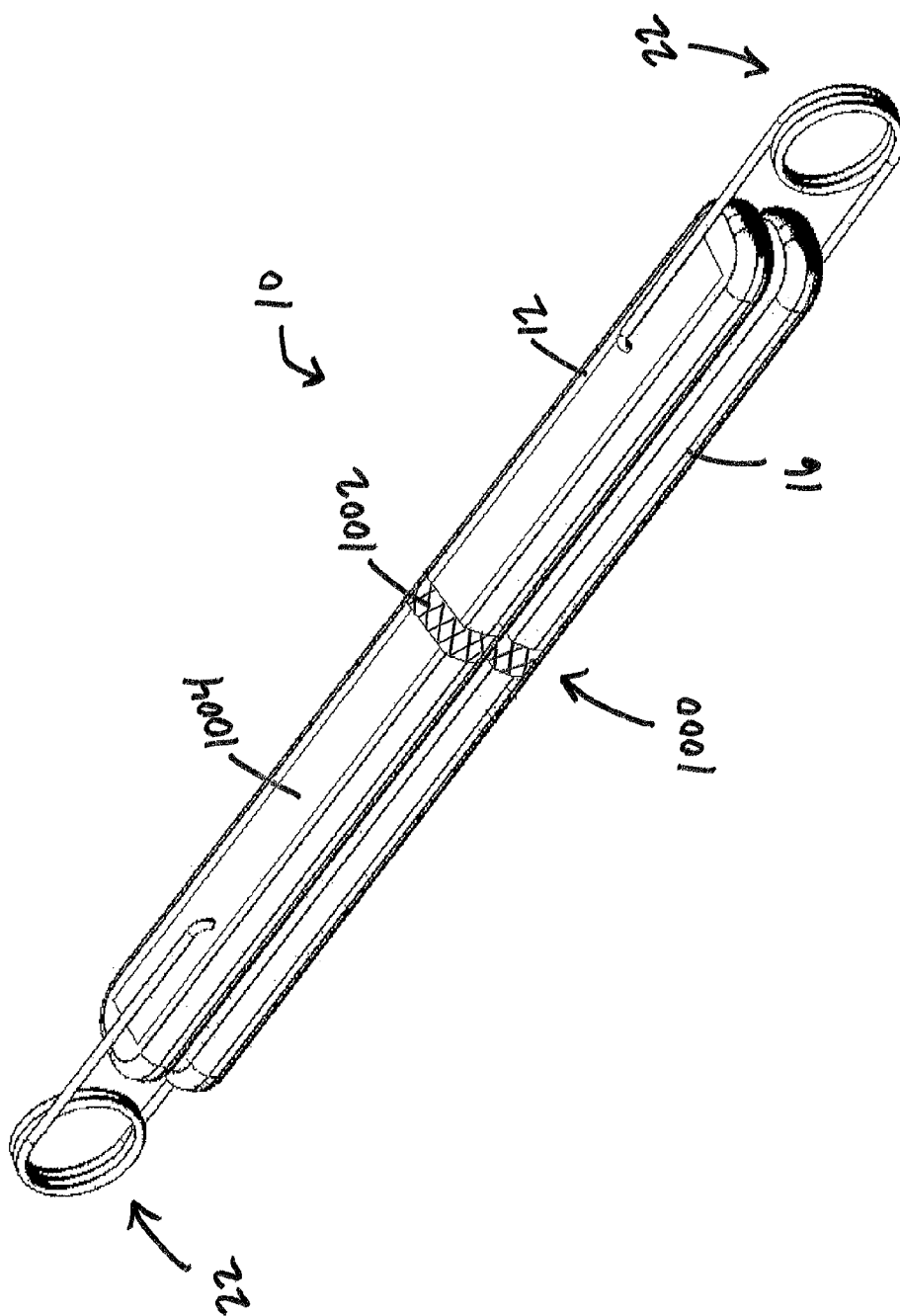


Fig. 10

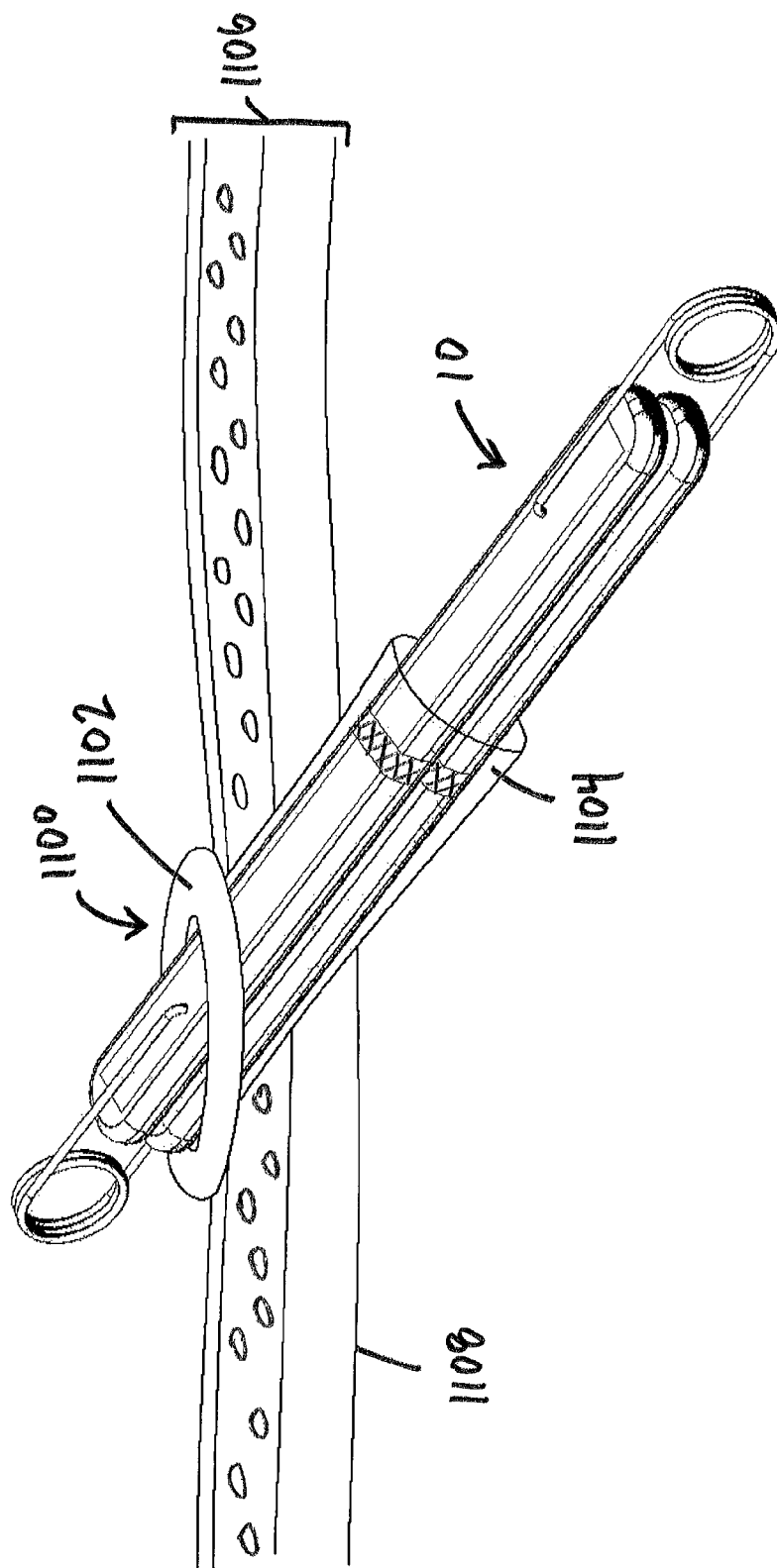
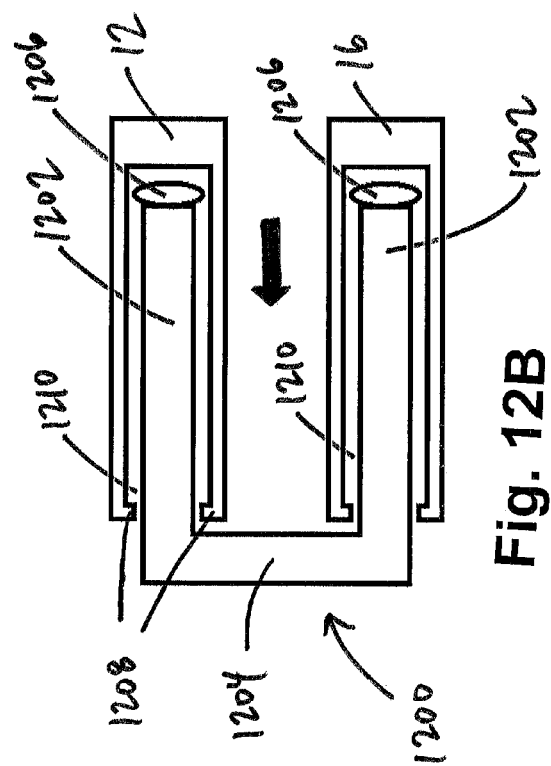
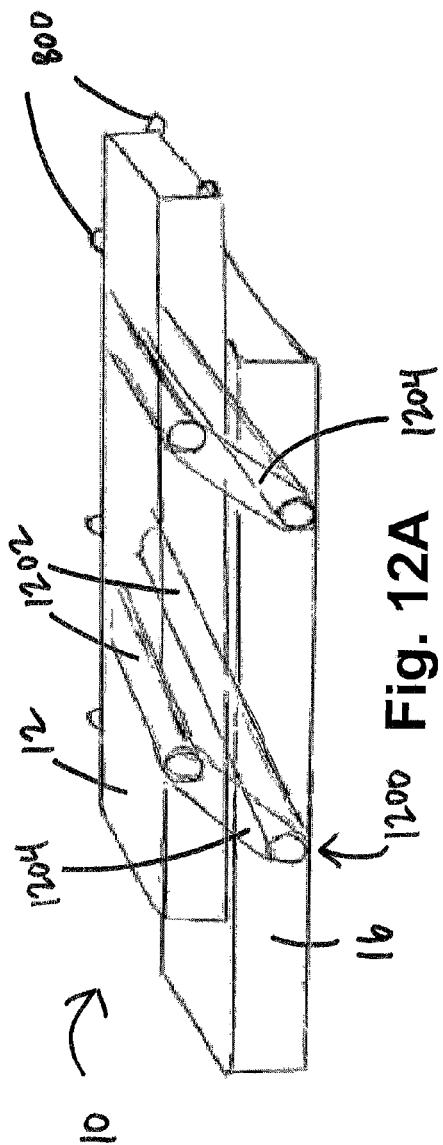
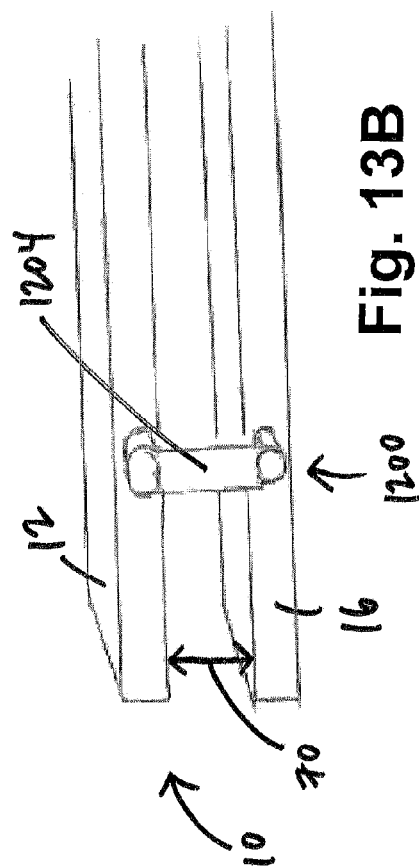
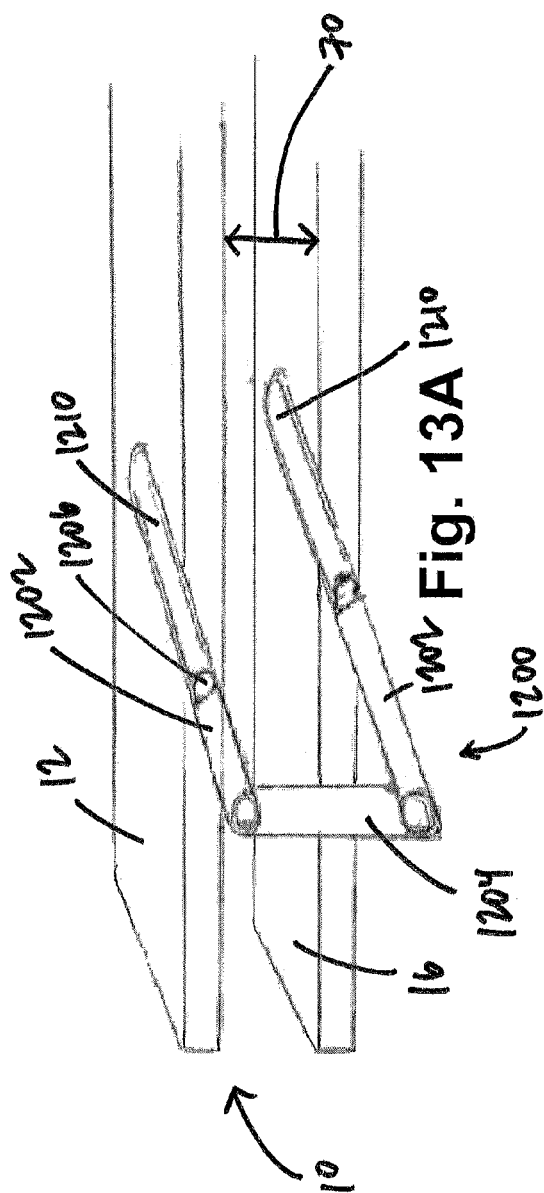
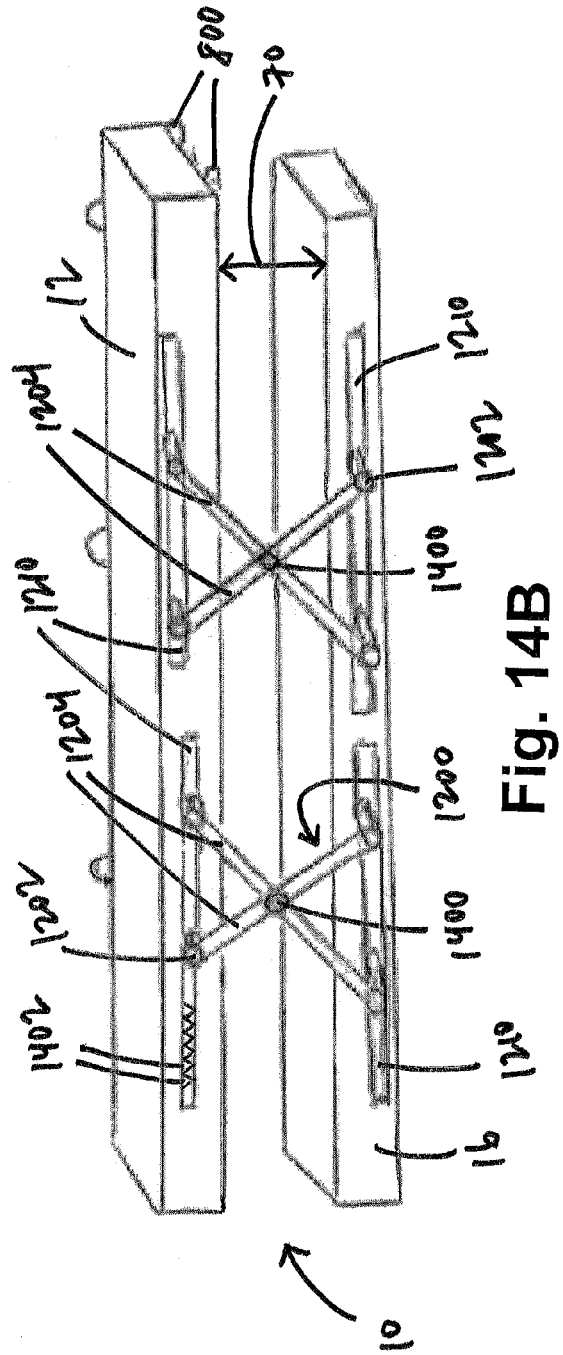
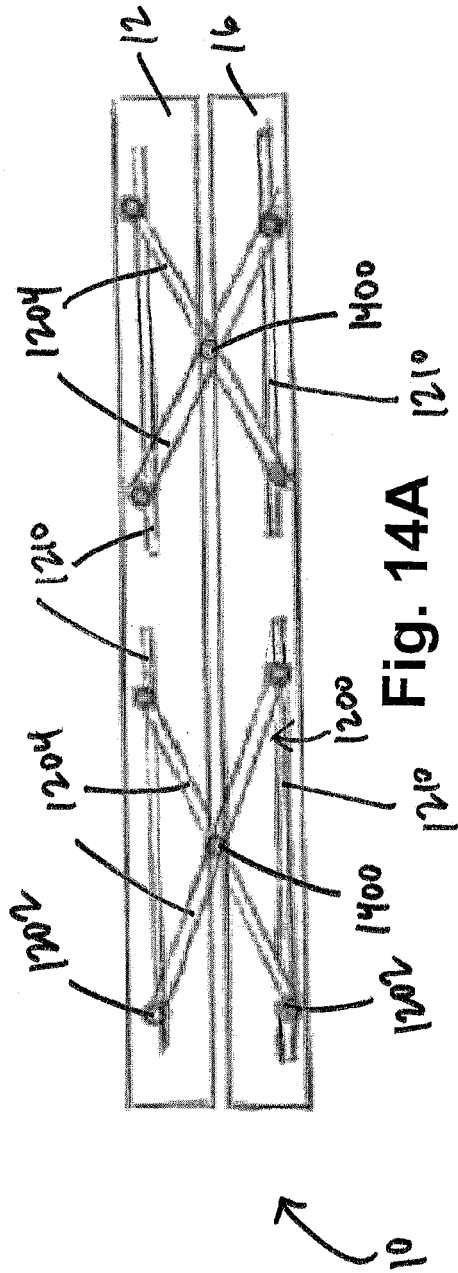
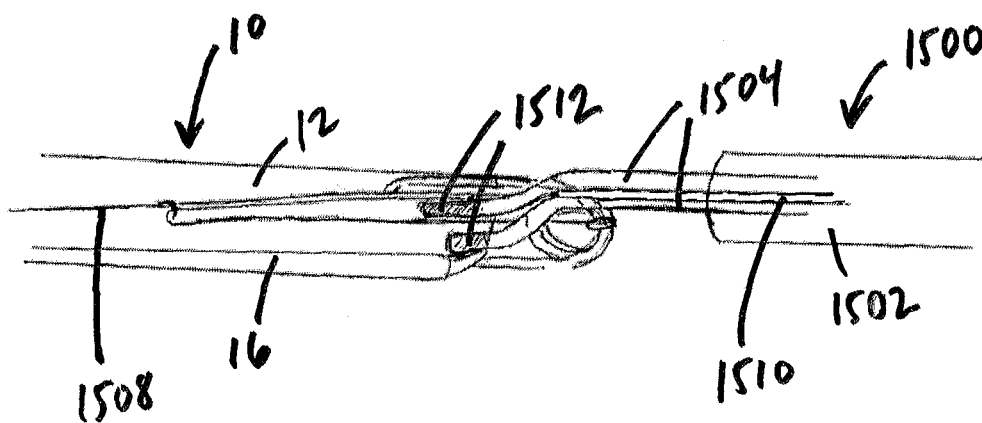
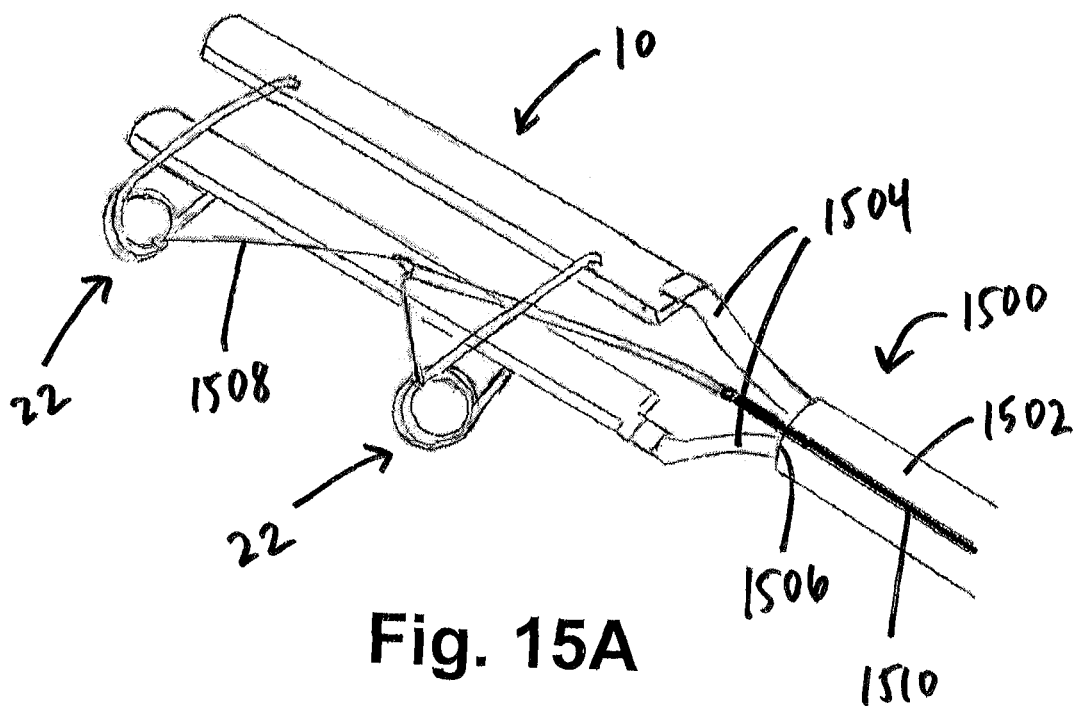


Fig. 11









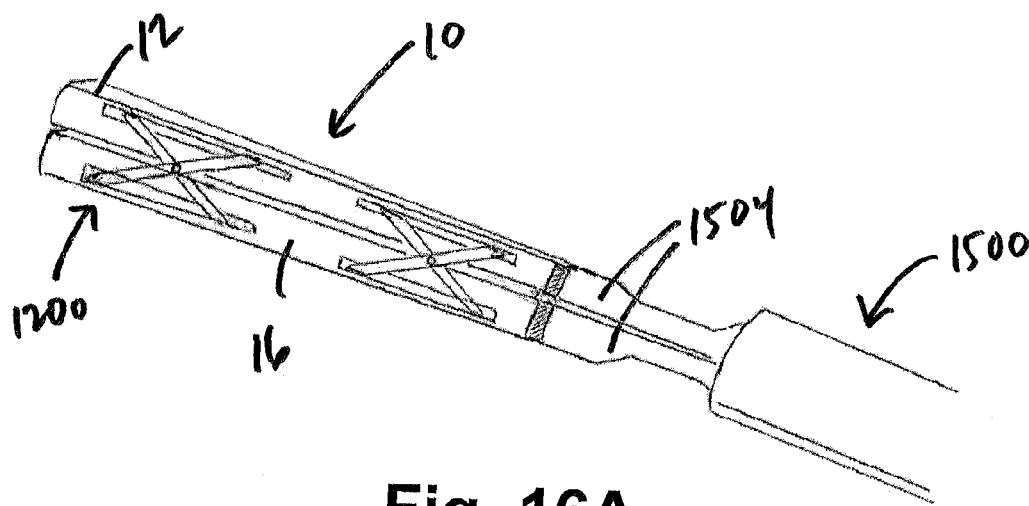


Fig. 16A

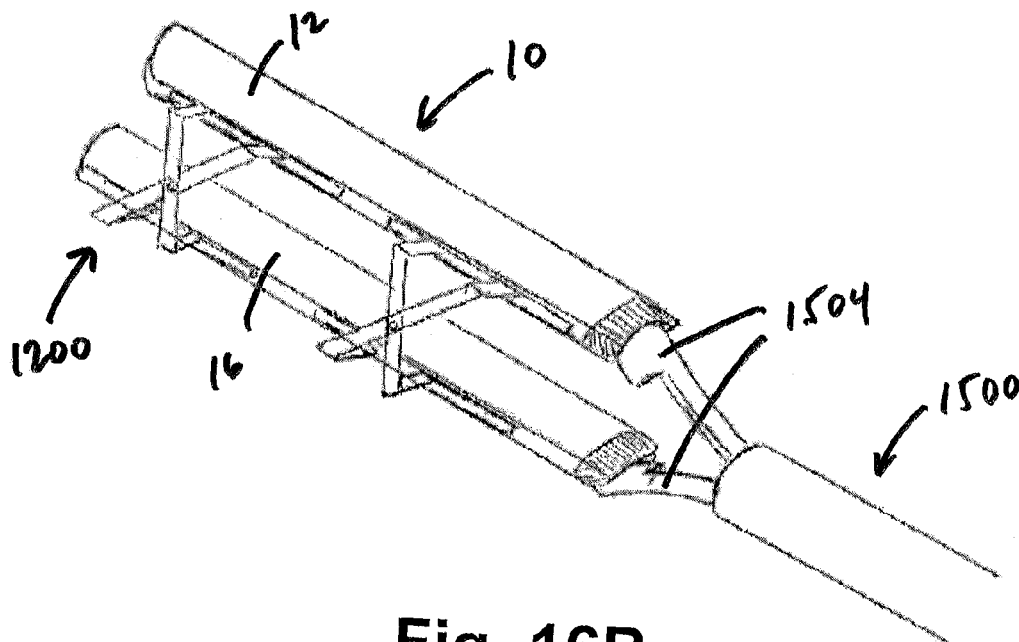
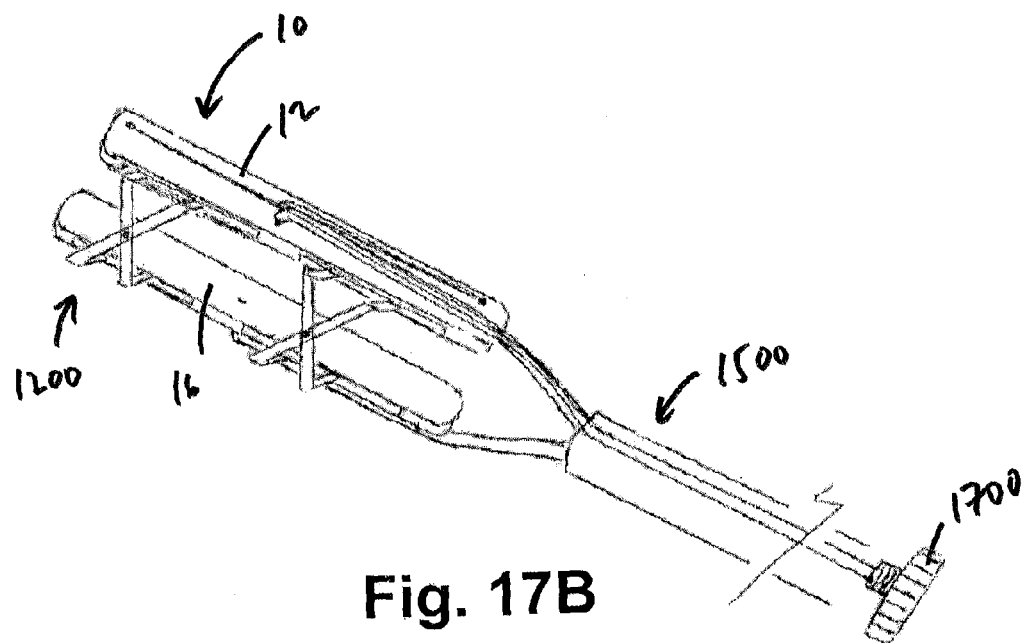
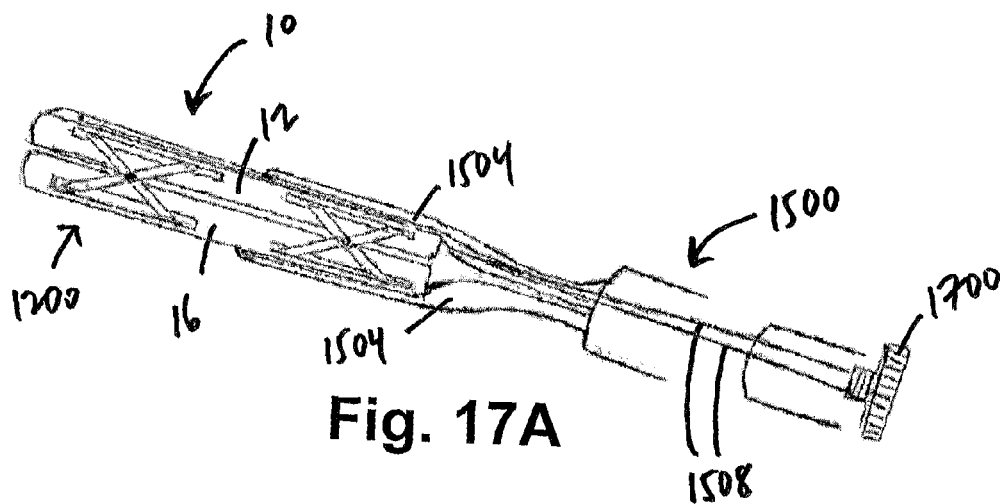


Fig. 16B



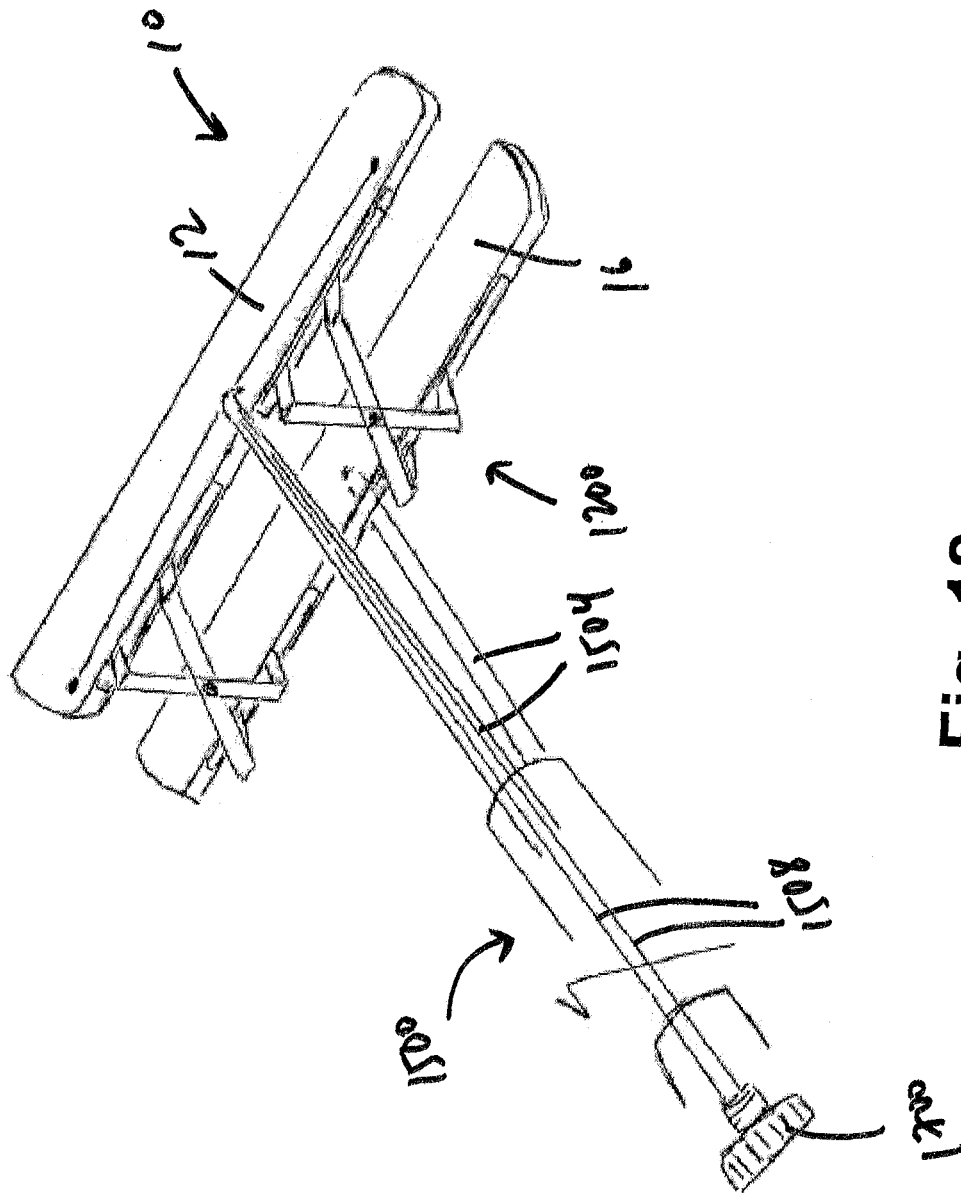
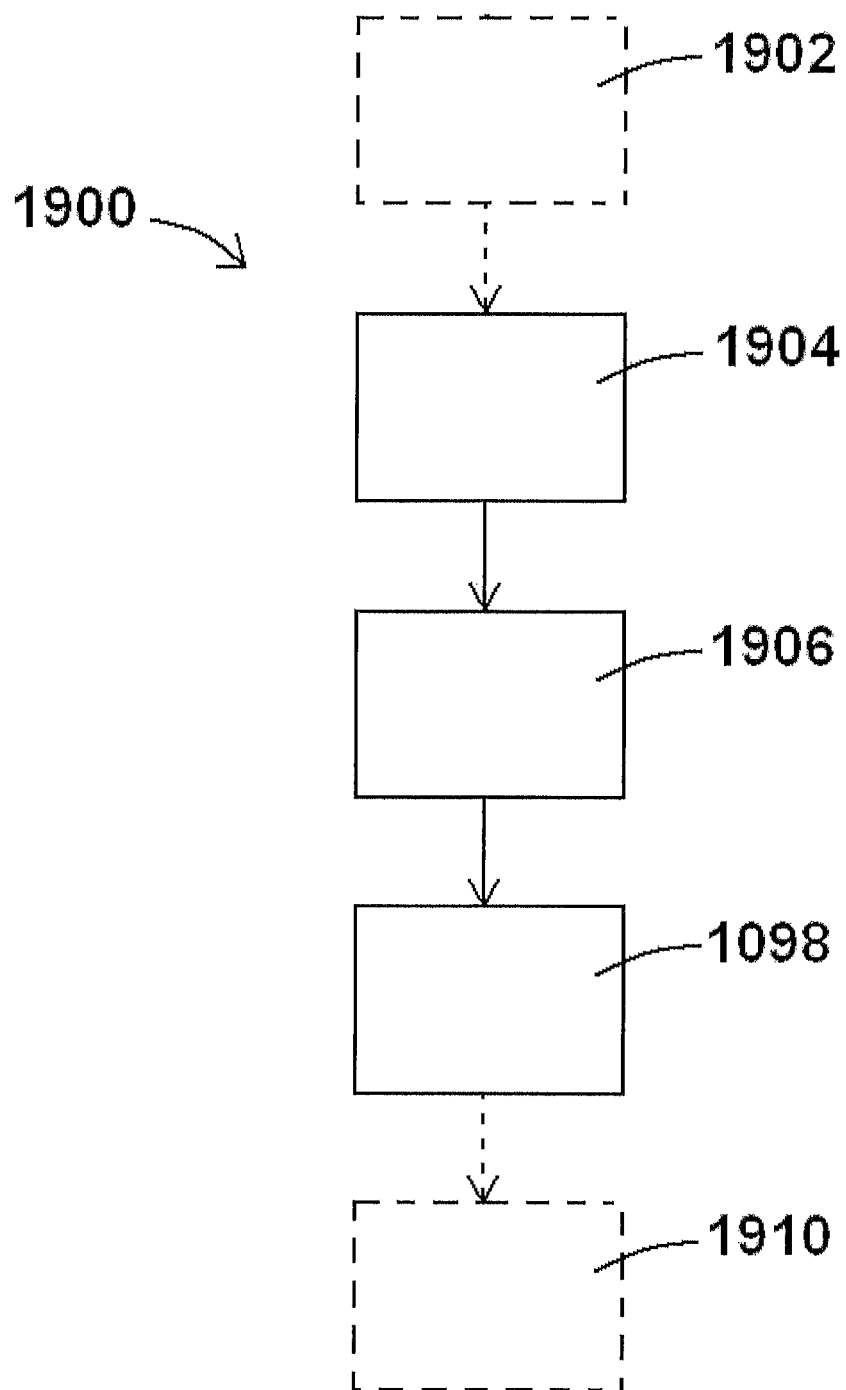


Fig. 18

**Fig. 19**

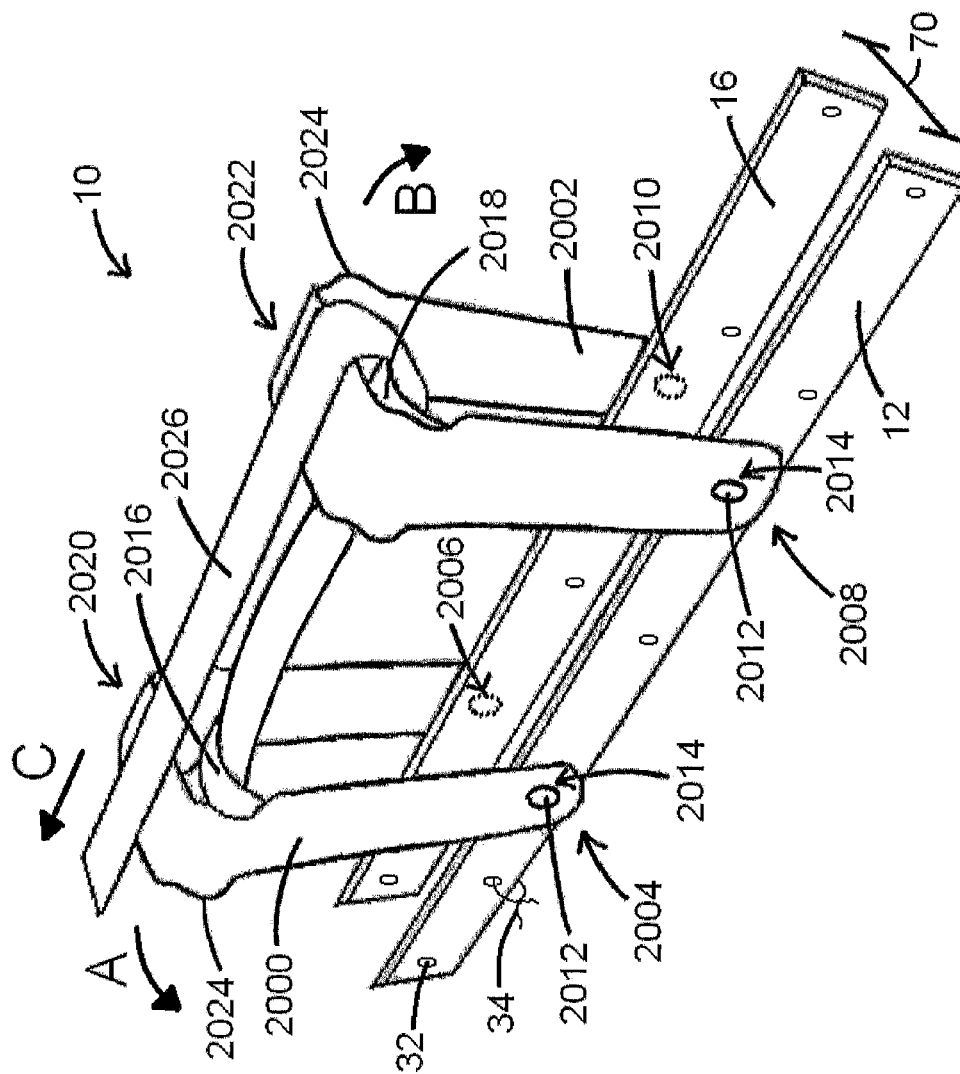


Fig. 20

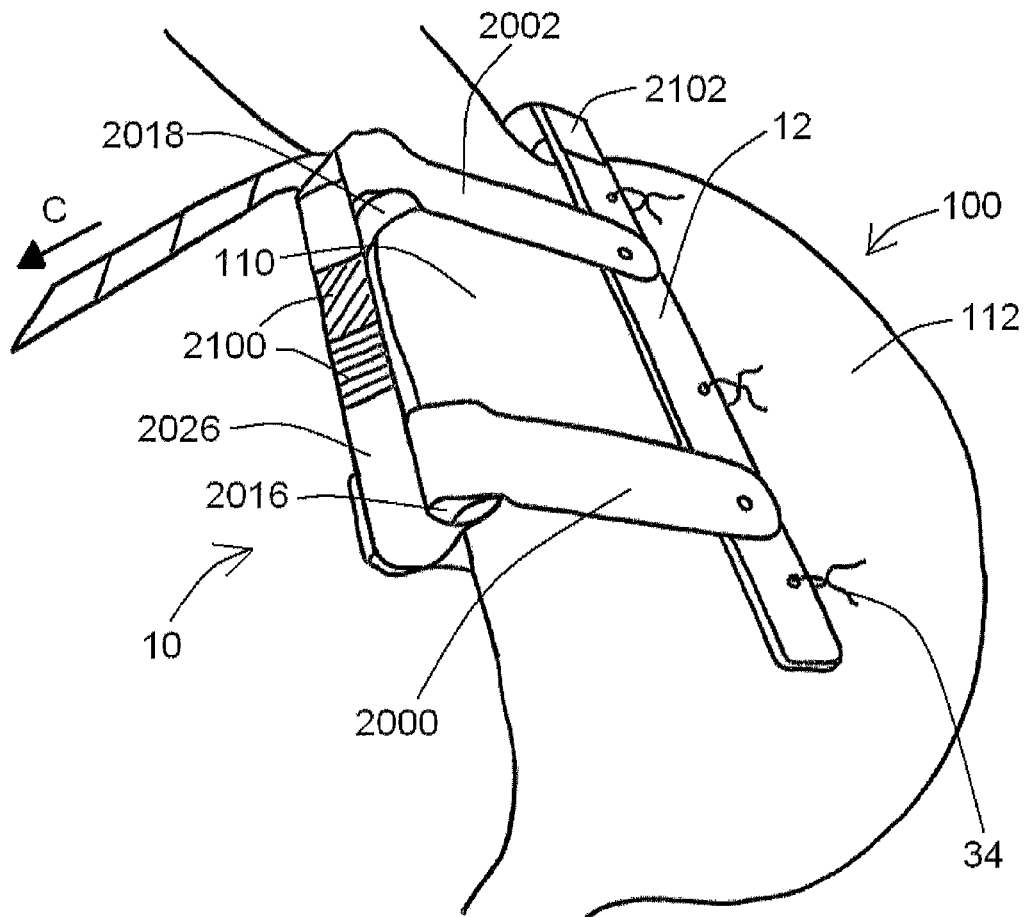
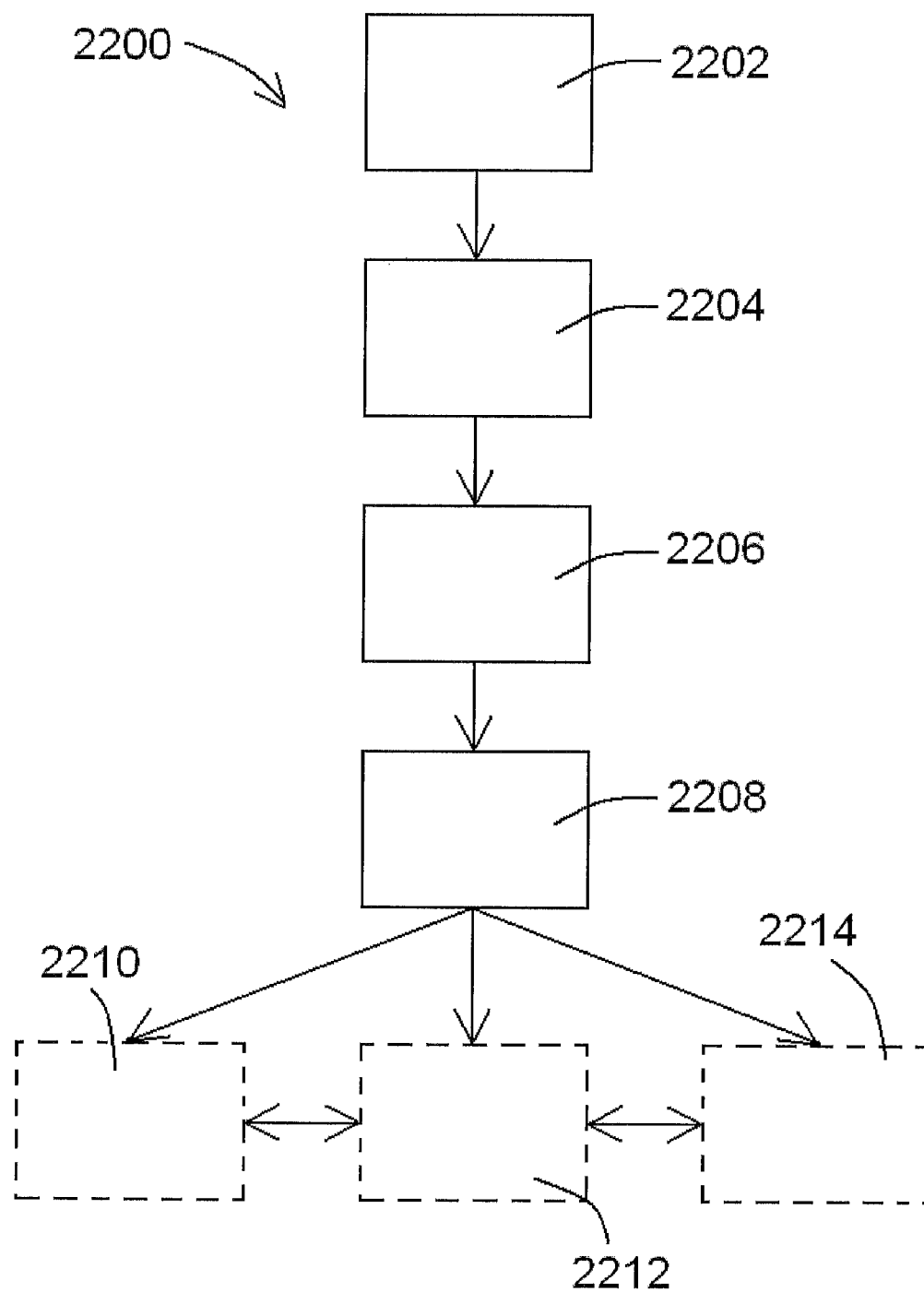


Fig. 21

**Fig. 22**

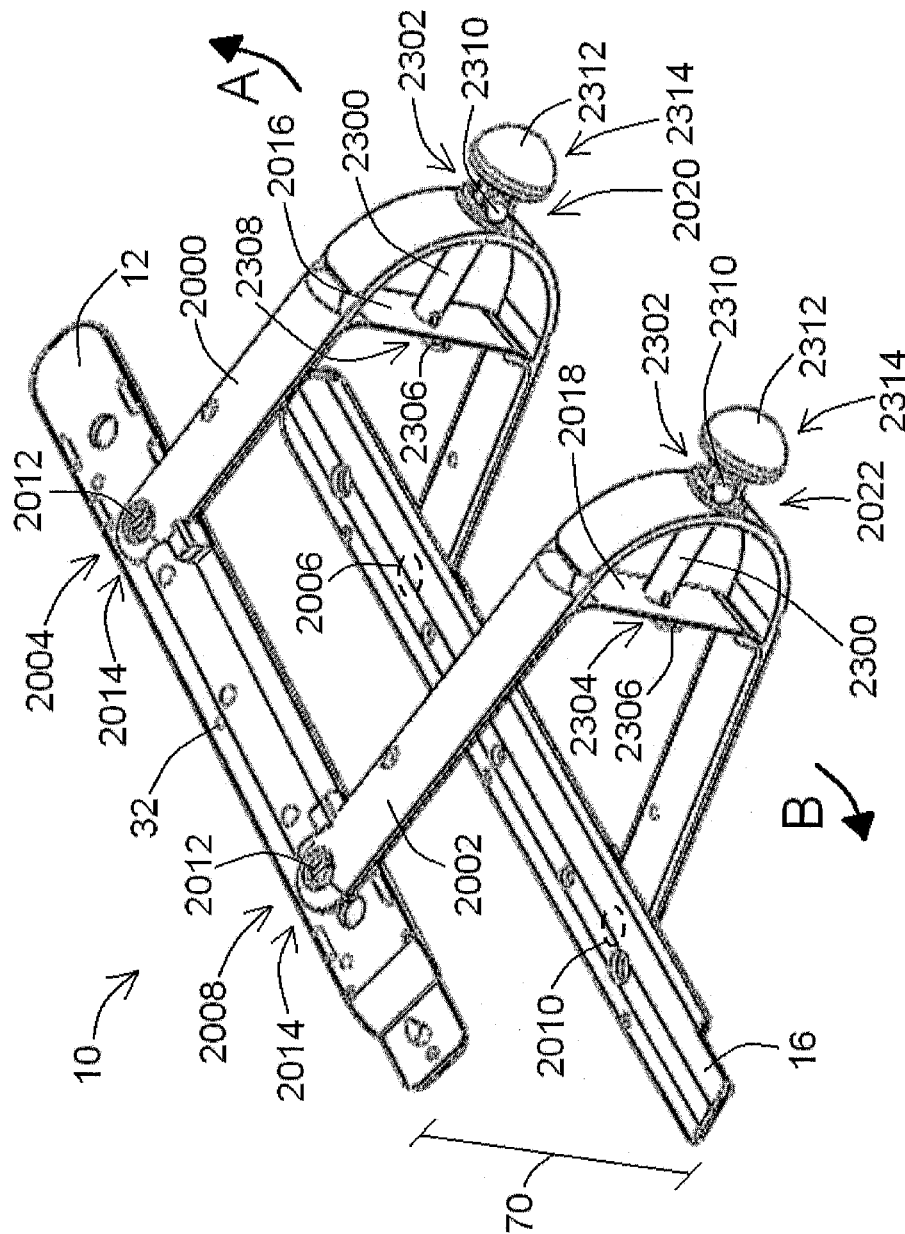


Fig. 23

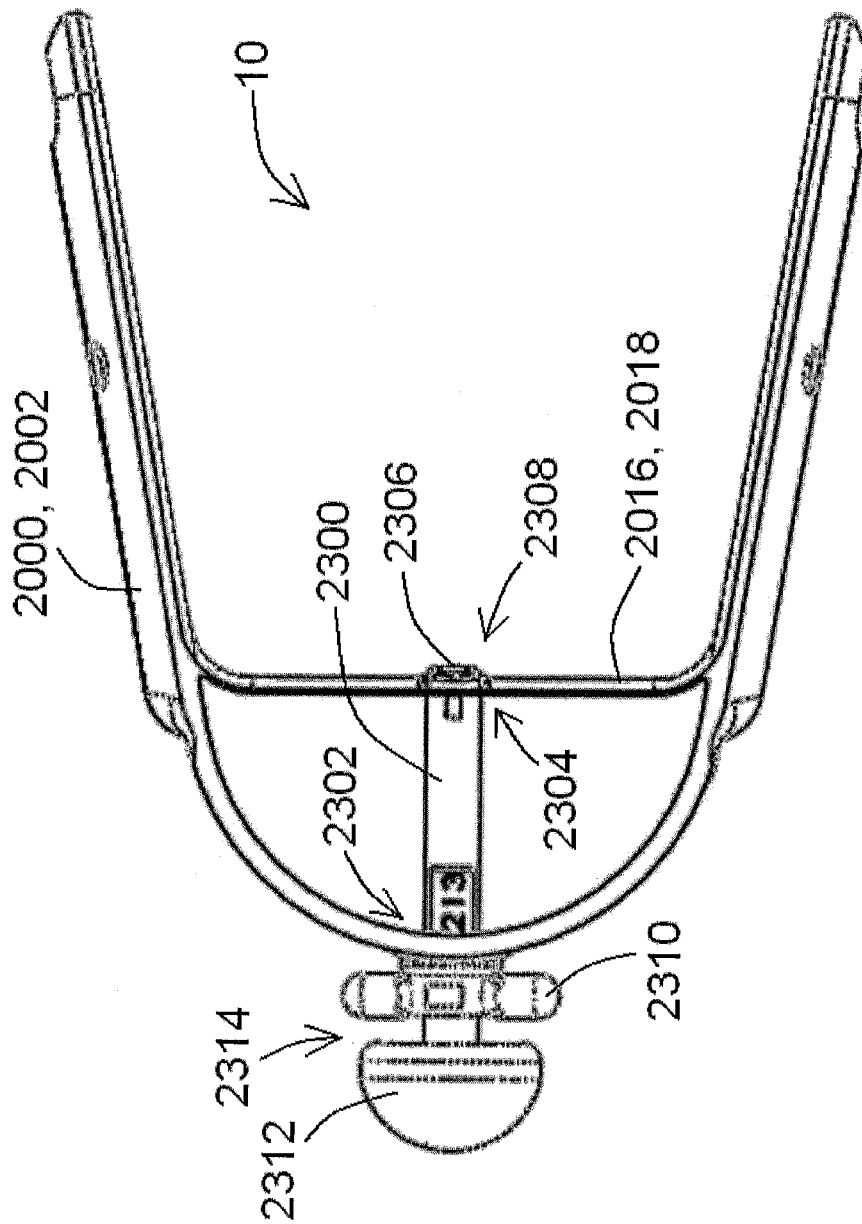


Fig. 24

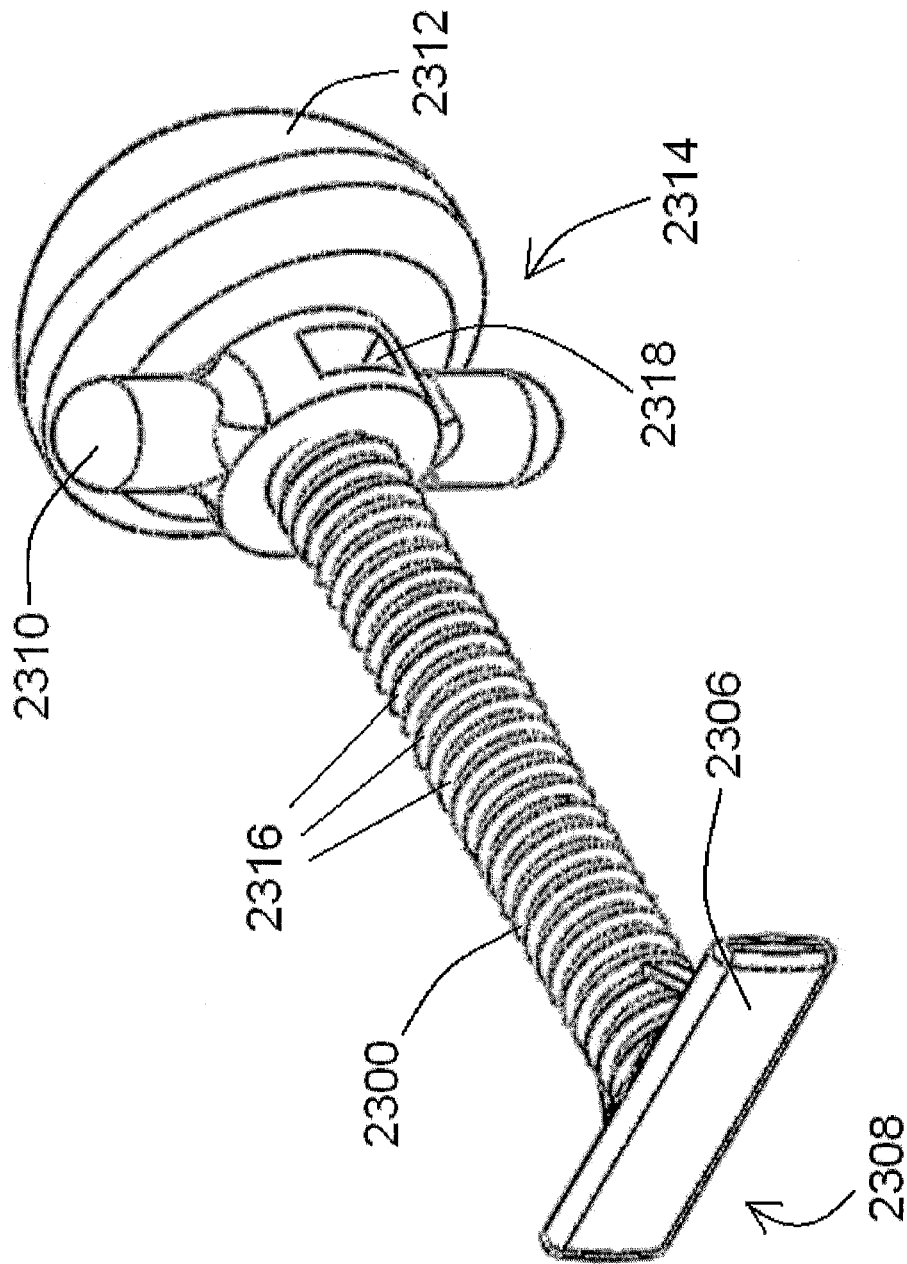


Fig. 25

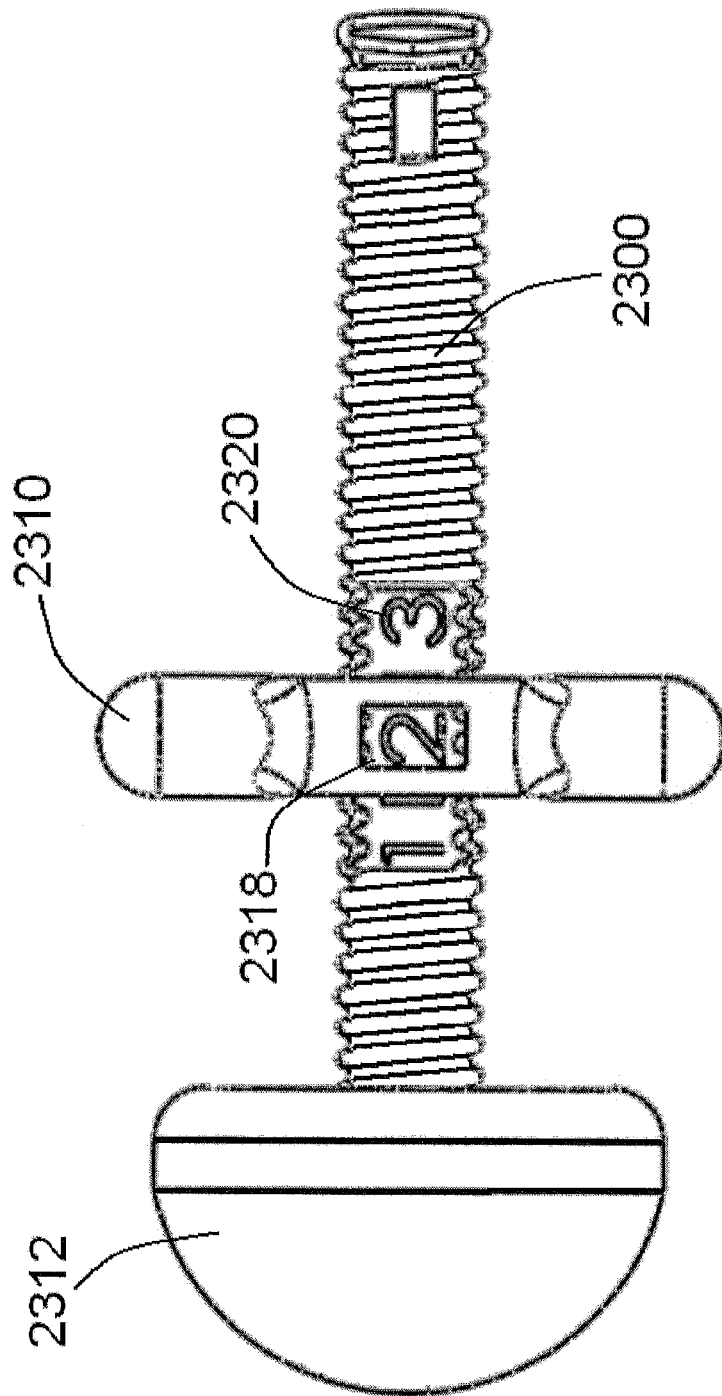


Fig. 26

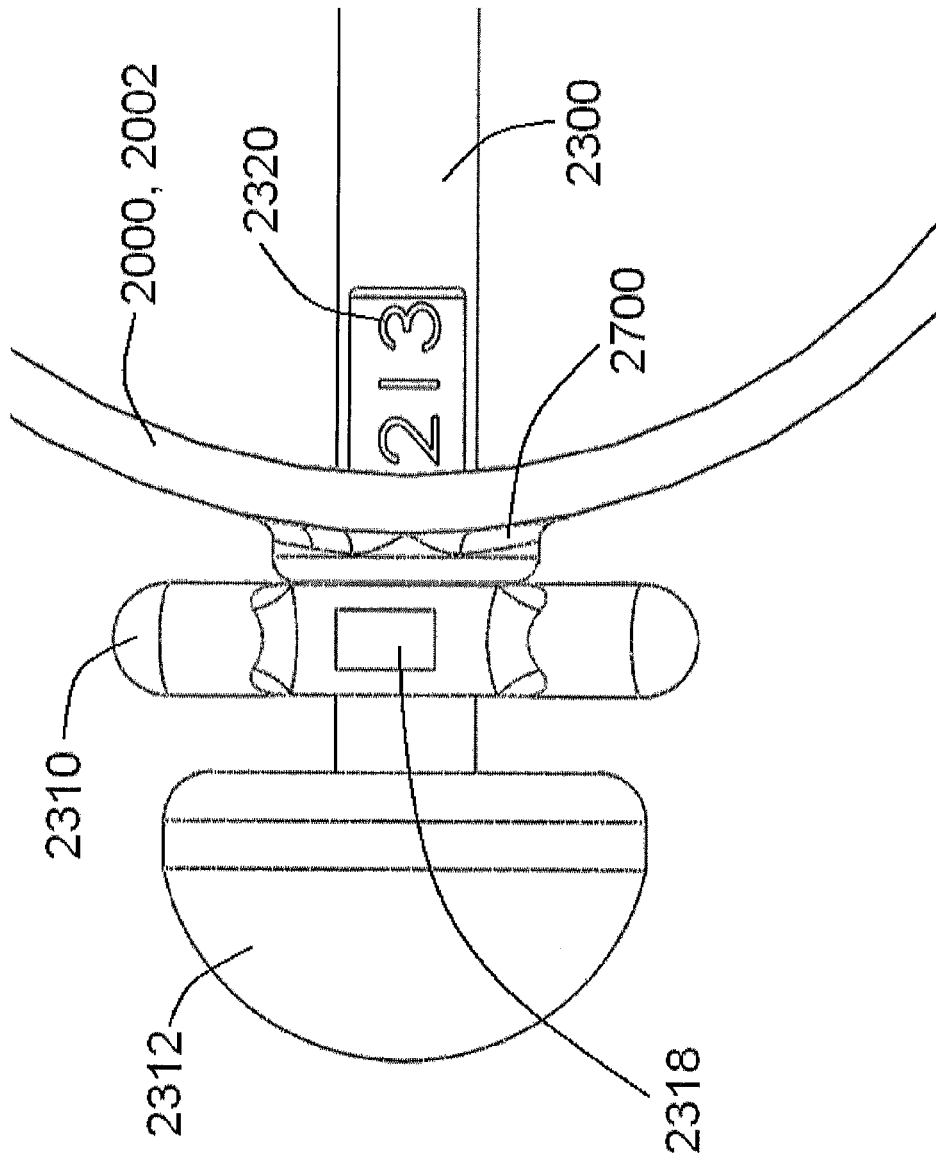


Fig. 27

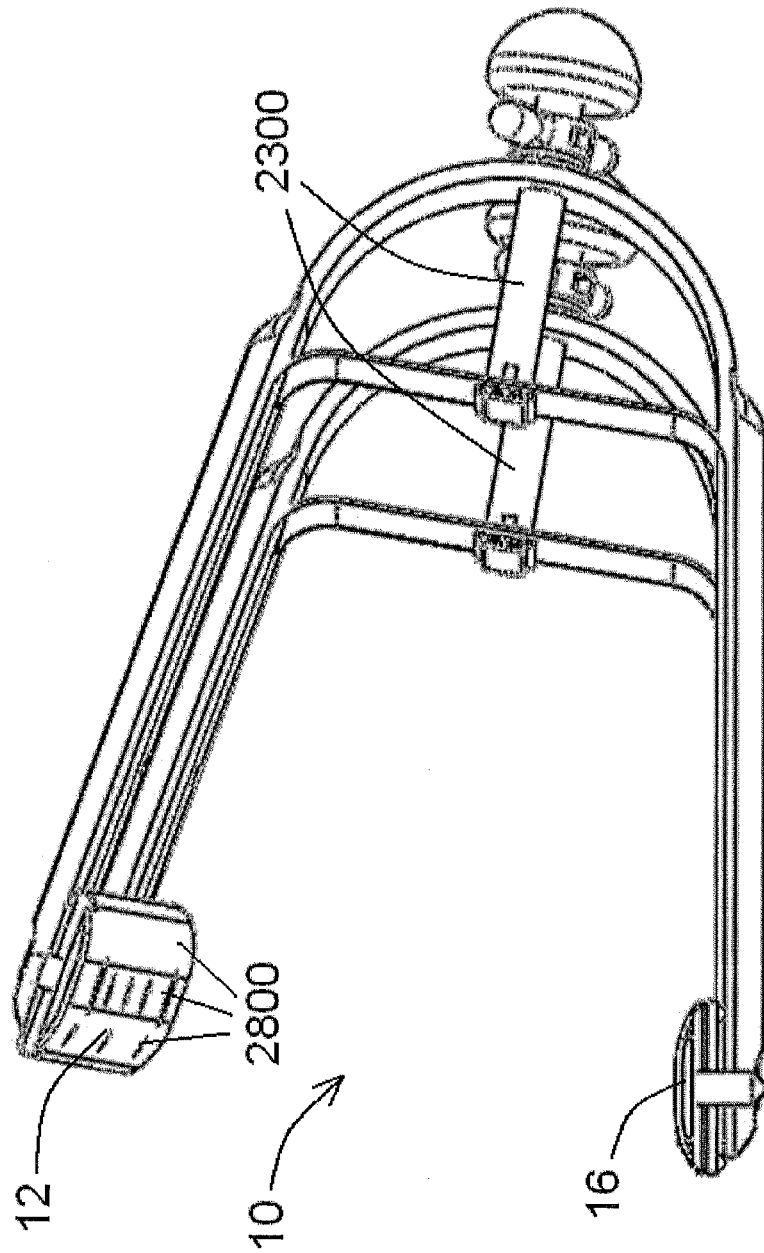
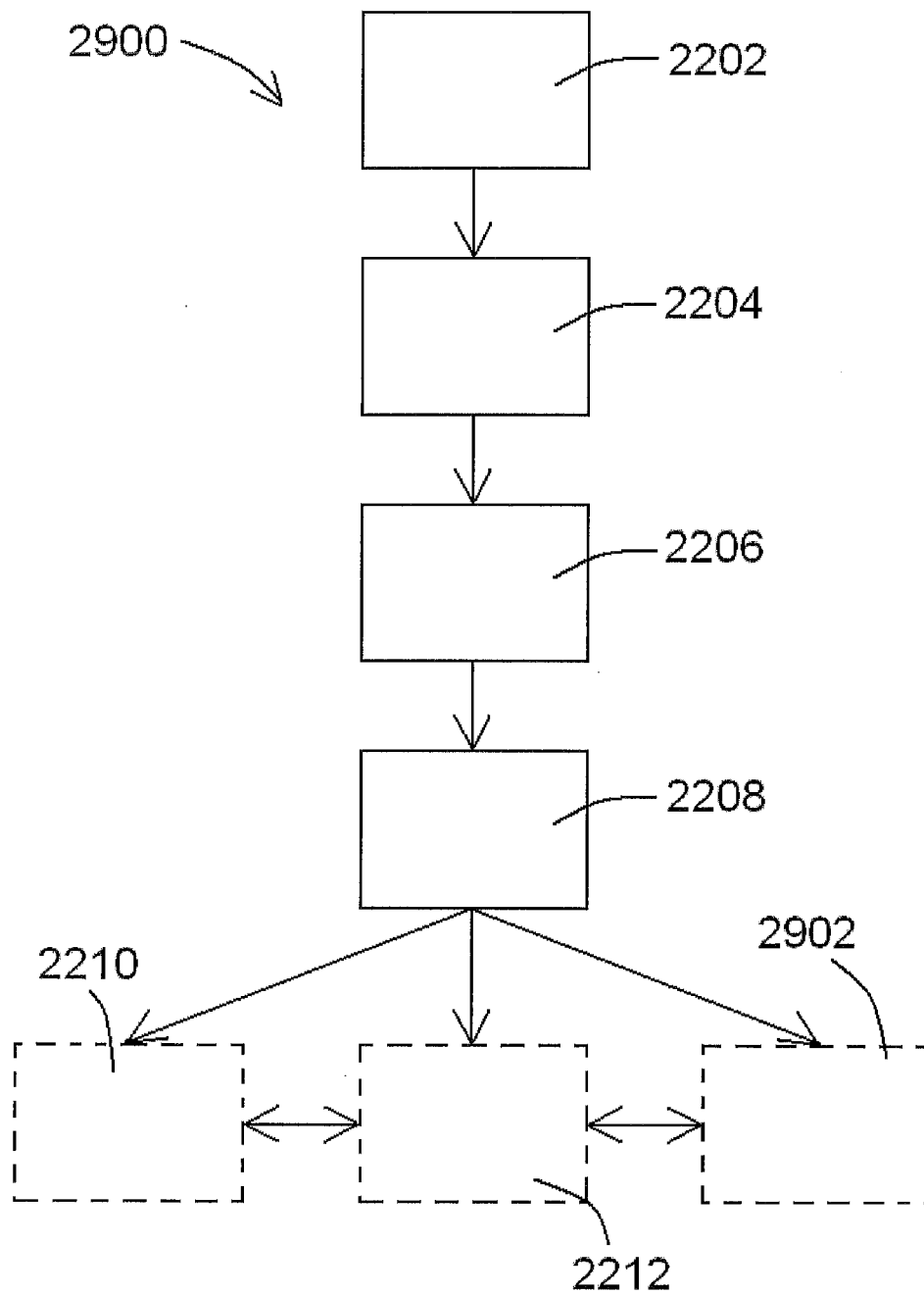


Fig. 28

**Fig. 29**

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TISSUE RESTORATION DEVICES, SYSTEMS, AND METHODS

PRIORITY

The present continuation-in-part patent application is related to, and claims the priority benefit of, U.S. patent application Ser. No. 12/546,139, filed Aug. 24, 2009, the contents of which are hereby incorporated by reference in their entirety into this disclosure.

BACKGROUND

Organ and tissue surgical restoration are clinical techniques that may be applied to numerous different body tissues, ranging from blood vessels to whole organs. Conventionally, such surgical techniques require the use of incisions, pins, staples and/or significant sutures in order to alter the tissue's anatomy. For example, surgical gastric restoration often employed to treat obesity and typically involves the reorganization of the digestive tract. Conventional examples of such procedures involve attempts to either 1) restrict food intake into the body via a restrictive bariatric procedure (a "Restrictive Procedure"), or 2) alter the anatomy of the small intestine or divert the peristalsis of a person's normal food intake past the small intestine to decrease caloric absorption via a malabsorptive bariatric procedure, which is commonly known as a gastric bypass (a "Malabsorptive Procedure"). It is also known to combine the two procedures such that both of the aforementioned techniques are employed jointly.

Malabsorptive Procedures entail an intestinal bypass that results in the exclusion of almost all of the small intestine from the digestive tract. In most Malabsorptive Procedures, a portion of the stomach or small intestine is removed from the digestive tract through a surgical procedure that requires cutting the digestive tissue and thereafter closing any holes or securing the newly formed anatomy with staples and/or sutures. Conversely, Restrictive Procedures generally involve the creation of a passageway extending from the upper portion of the stomach to the lower portion of the stomach in order to decrease the size of the organ and thus prevent the stomach from storing large amounts of food. Conventional Restrictive Procedures rely on the banding, suturing and/or stapling of the stomach to create a small pouch on the superior portion of the stomach near the gastroesophageal junction.

Combined operations consisting of Malabsorptive and Restrictive Procedures are the most common bariatric procedures performed today. An example of a combined procedure is the Extended (Distal) Roux-en-Y Gastric Bypass in which a stapling creates a small stomach pouch (approximately 15 to 20 cc) completely separated from the remainder of the stomach. In addition, the small intestine is divided just beyond the duodenum (the hollow tube connecting the stomach to the jejunum), re-arranged into a Y-configuration, and sutured to the small upper stomach pouch to enable the outflow of food therefrom through the newly formed "Roux limb,"

Accordingly, most digestive tract restoration procedures require that the stomach and/or tissue of the intestine is cut and thereafter sutured or stapled back together. As the digestive tract contains numerous enzymes, strong acids and multiple species of bacteria that assist with digestion, the perforation of an organ and/or tissue thereof is particularly problematic due to the likelihood of leakage therefrom and/or increased risk of serious infection. As such, conventional gastric surgical restoration procedures have high rates of post-operative complications that may require prolonged

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hospitalization and even additional operations, and are often irreversible and/or permanently affect the restored tissue and/or organ. Accordingly, a need exists for safe and effective devices and methods for restoring organs and tissue that are reversible and do not require cutting or penetrating the underlying tissue with significant sutures, staples and/or pins.

It will be appreciated that the foregoing examples are only provided as examples and that there are numerous other indications where intervention is necessary either to restore the underlying organ or tissue and/or to provide support thereto.

BRIEF SUMMARY

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, the device comprises a first engaging component and a second engaging component, each component configured for laparoscopic insertion into a body cavity, a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, and a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend. In another embodiment, the first swivel arm and the second swivel arm are capable of moving between a first position that is substantially parallel with the first engaging component and the second engaging component and a second position that is substantially perpendicular with the first engaging component and the second engaging component. In yet another embodiment, the first engaging component and the second engaging component are configured to engage a targeted tissue therebetween when the first swivel arm and the second swivel arm are in a configuration relatively perpendicular to the first engaging component and the second engaging component. In an additional embodiment, the first engaging component and the second engaging component each define one or more studs sized and shaped to permit the first swivel arm and the second swivel arm to be coupled thereto.

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, the first swivel arm and the second swivel arm further define one or more curvatures, whereby the first bend, the second bend, and the one or more curvatures define a native U-shaped configuration. In an additional embodiment, the device further comprises a tape positioned around at least part of the device so that the tape engages the first interconnection arm and the second interconnection arm. In another embodiment, the tape is capable of decreasing an interior space defined between the first engaging component and the second engaging component when the tape applies a force to the first interconnection arm and the second interconnection arm. In yet another embodiment, the tape further comprises one or more detectable portions positioned/imprinted thereon. In various embodiments, the one or more detectable portions are radio-opaque.

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, the device further comprises a cover flap, the cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging

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component when initially coupled to the second engaging component. In another embodiment, the at least one of the first engaging component and the second engaging component define one or more suture apertures therethrough. In yet another embodiment, the first engaging component and the second engaging component each comprise a proximal end, a distal end, and a body extending therebetween, wherein the body of the first engaging component is configured to conform to a first targeted tissue surface, and the body of the second engaging component is configured to conform to a second targeted tissue surface.

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, the device further comprises a first adjustment rod and a second adjustment rod, each of the first adjustment rod and the second adjustment rod comprising a distal end, a proximal end, and threading therebetween, and a dial rotatably coupled thereto at or near the threading. In another embodiment, the first adjustment rod is coupled to the first swivel arm and the first interconnection arm, and wherein the second adjustment rod is coupled to the second swivel arm and the second interconnection arm. In yet another embodiment, rotation of the dial on either or both of the first adjustment rod and/or the second adjustment rod facilitates movement of the first engaging component and the second engaging component. In an additional embodiment, rotation of the dial in a first direction causes the first engaging component and the second engaging component to move toward one another, and wherein rotation of the dial in a second direction causes the first engaging component and the second engaging component to move away from one another.

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, each of the first adjustment rod and the second adjustment rod further comprises a bar coupled thereto at or near the distal ends of said rods. In an additional embodiment the bar coupled to the first adjustment rod is positioned distal to the first interconnection arm, and wherein the bar coupled to the second adjustment rod is positioned distal to the second interconnection arm, and wherein rotation of the dial on either or both of the first adjustment rod and/or the second adjustment rod facilitates movement of the first engaging component and the second engaging component. In yet an additional embodiment each of the first adjustment rod and the second adjustment rod further comprises a cap coupled thereto at or near the proximal ends of said rods, said caps configured to prevent said dials from disengaging said adjustment rods. In another embodiment, said dials define a dial aperture therethrough, the dial configured to permit an indicia present upon said adjustment rods to be viewed therethrough. In yet another embodiment, each of the first engaging component and the second engaging component define one or more facets along at least part of a length of said engaging components, said facets providing a generally arcuate profile of said engaging components.

In at least one exemplary embodiment of an implantable restraining device of the present disclosure, each of the first engaging component and the second engaging component comprise a configuration chosen from a straight bar configuration, a curved configuration, or a circular configuration. In an additional embodiment, the first engaging component and the second engaging component are flexible or semi-flexible. In various embodiments, the first engaging component, the second engaging component, the first swivel arm, and the second swivel arm each comprise a material suitable to resist corrosion selected from the group consisting of polyurethane, polyethylene, polytetrafluoroethylene, nitinol, silastic, titanium, and stainless steel. In various embodiments, the first

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engaging component, the second engaging component, the first swivel arm, and the second swivel arm comprise a resorbable material selected from the group consisting of polyglycolide (PGA), polylactide (PLA), L-lactide (LPLA), poly(DL-lactide) (DLPLA), poly(ϵ -caprolactone) (PCL), poly(dioxanone) (PDO), polyglycolide-trimethylene carbonate (PGA-TMC), or poly(D,L-lactide-co-glycolide) (DLPLG).

In at least one exemplary embodiment of implantable restraining device of the present disclosure, the device comprises a first engaging component defining a first bend and a second engaging component defining a second bend, the first and second engaging components configured for laparoscopic insertion into a body cavity, a first swivel arm and a second swivel arm, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend, a tape positioned around at least part of the device so that the tape engages the first interconnection arm and the second interconnection arm, and a cover flap, the cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging component when initially coupled to the second engaging component, wherein the first swivel arm and the second swivel arm are capable of moving between a first position that is substantially parallel with the first engaging component and the second engaging component and a second position that is substantially perpendicular with the first engaging component and the second engaging component.

In at least one exemplary embodiment of implantable restraining device of the present disclosure, the device comprises a first engaging component and a second engaging component, each component configured for laparoscopic insertion into a body cavity, a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend, a first adjustment rod and a second adjustment rod, each of the first adjustment rod and the second adjustment rod comprising a distal end, a proximal end, and threading therebetween, a dial rotatably coupled thereto at or near the threading, and a bar coupled thereto at or near the distal ends of said adjustment rods, wherein the first adjustment rod is coupled to the first swivel arm and the first interconnection arm, and wherein the second adjustment rod is coupled to the second swivel arm and the second interconnection arm, wherein rotation of the dial in a first direction causes the first engaging component and the second engaging component to move toward one another, and wherein rotation of the dial in a second direction causes the first engaging component and the second engaging component to move away from one another, and wherein the first swivel arm and the second

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swivel arm are capable of moving between a first position that is substantially parallel with the first engaging component and the second engaging component and a second position that is substantially perpendicular with the first engaging component and the second engaging component.

In at least one exemplary embodiment of a method for restoring a targeted tissue of the present disclosure, the method comprises the steps of inserting an implantable restraining device into a body cavity of a mammalian body, the implantable restraining device comprising a first engaging component defining a first bend and a second engaging component defining a second bend, the first and second engaging components configured for laparoscopic insertion into a body cavity, a first swivel arm and a second swivel arm, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, and a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend, advancing the implantable restraining device to a location within the mammalian body adjacent to a targeted tissue, swiveling the first swivel arm and the second swivel arm so that the first swivel arm and the second swivel arm are substantially perpendicular to the first engaging component and the second engaging component, and positioning the first engaging component and the second engaging component over the targeted tissue such that at least a portion of the targeted tissue is positioned therebetween, wherein when the targeted tissue expands in a direction between the first engaging component and the second engaging component, the targeted tissue exerts a force upon the first engaging component and the second engaging component. In another embodiment, the method further comprises the step of securing one or more sutures to connect the first engaging component and/or the second engaging component to the targeted tissue. In yet another embodiment, the method further comprises the step of securing a cover flap coupled to either the first engaging component or the second engaging component so that the cover flap capable is either further coupled to the second engaging component when initially coupled to the first engaging component or further coupled to the first engaging component when initially coupled to the second engaging component. In an additional embodiment, the method further comprises the step of adjusting a tape positioned around at least part of the device whereby the tape engages the first interconnection arm and the second interconnection arm, wherein adjustment of the tape decreases an interior space defined between the first engaging component and the second engaging component when the tape applies a force to the first interconnection arm and the second interconnection arm, and wherein adjustment of the tape increases an interior space defined between the first engaging component and the second engaging component when the force is removed or reduced. In various embodiments, the targeted tissue is a stomach, and wherein expansion of the stomach, with said device positioned thereon, functionally divides the stomach into a first stomach portion and a second stomach portion. In at least another embodiment, the implantable restraining device further comprises a first adjustment rod and a second adjustment rod, each of the first adjustment rod and the second adjustment rod having a dial rotatably coupled thereto, wherein the first adjustment rod is coupled to the first swivel arm and the first interconnection arm, and wherein the second adjustment

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rod is coupled to the second swivel arm and the second interconnection arm, and wherein the method further comprises the step, of rotating said dials to adjust an interior space between the first engaging component and the second engaging component.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1 and 2 show perspective views of at least one embodiment of a restraining device for restoring and/or supporting a tissue or organ of the present disclosure;

FIG. 3A shows a top view of at least one embodiment of a restraining device for restoring and/or supporting a tissue or organ according to the present disclosure;

FIGS. 3B through 3E show various embodiments of engaging components according to the present disclosure;

FIG. 4A shows a side view of the restraining device of FIGS. 1A and 1B applied in a longitudinal fashion to a stomach according to the present disclosure;

FIG. 4B shows a cross-sectional view of a restored stomach along line A-A of FIG. 4;

FIGS. 5A through 5D show various embodiments of restraining devices positioned about a stomach according to the present disclosure;

FIG. 6 shows a flow chart of a method for laparoscopically delivering embodiments of the restraining device disclosed herein to a targeted tissue according to the present disclosure;

FIGS. 7A, 7B, and 8 show various embodiments of a restraining device comprising one or more struts according to the present disclosure;

FIG. 9 shows an exemplary embodiment of a restraining device comprising two springs and a mesh curtain according to the present disclosure;

FIG. 10 shows an exemplary embodiment of a restraining device comprising a pliable junction according to the present disclosure;

FIG. 11 shows a restraining device positioned within a laparoscopic port according to the present disclosure;

FIGS. 12A, 12B, 13A, and 13B show various embodiments of a restraining device comprising a coupler according to the present disclosure;

FIGS. 14A and 14B show exemplary embodiments of a restraining device comprising a coupler and a coupler pivot according to the present disclosure;

FIGS. 15A and 15B show exemplary embodiments of an apparatus for delivering a restraining device according to the present disclosure;

FIGS. 16A and 16B show exemplary embodiments of an apparatus for delivering a restraining device comprising a coupler according to the present disclosure;

FIGS. 17A and 17B show additional exemplary embodiments of an apparatus for delivering a restraining device according to the present disclosure;

FIG. 18 shows an exemplary embodiment of an apparatus perpendicularly engaging a restraining according to the present disclosure;

FIG. 19 shows a flow chart of a method for laparoscopically removing embodiments of the restraining device disclosed herein from a targeted tissue according to the present disclosure;

FIG. 20 shows a perspective view of at least another embodiment of a restraining device for restoring and/or supporting a tissue or organ of the present disclosure;

FIG. 21 shows an exemplary embodiment of a restraining device positioned about a stomach according to the present disclosure;

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FIG. 22 shows steps of an exemplary method of inserting, delivering, and placing a restraining device about a targeted tissue according to the present disclosure;

FIG. 23 shows a perspective view of at least another embodiment of a restraining device for restoring and/or supporting a tissue or organ of the present disclosure;

FIG. 24 shows a side view of the embodiment of a restraining device shown in FIG. 23;

FIG. 25 shows a perspective view of an embodiment of an adjustment rod of the present disclosure;

FIG. 26 shows a side view of the embodiment of an adjustment rod shown in FIG. 25;

FIG. 27 shows a side view of an embodiment of an adjustment rod coupled to at least another component of an exemplary restraining device of the present disclosure;

FIG. 28 shows another perspective view of the embodiment of a restraining device shown in FIG. 23; and

FIG. 29 shows steps of another exemplary method of inserting, delivering, and placing a restraining device about a targeted tissue according to the present disclosure.

DETAILED DESCRIPTION

For the purposes of promoting an understanding of the principles of the present disclosure, reference will now be made to the embodiments illustrated in the drawings, and specific language will be used to describe the same. It will nevertheless be understood that no limitation of the scope of this disclosure is thereby intended.

FIGS. 1 and 2 show perspective views of at least one embodiment of a restraining device 10 for restoring a tissue or organ. In at least one embodiment of the restraining device 10, the restraining device 10 comprises an implantable device and does not require substantial sutures, staples or pins to be secured on the targeted tissue or organ of interest. Further, the restraining device 10 described herein is configured to loosely, albeit securely, engage to the targeted tissue such that the underlying tissue is either uncompressed or only loosely compressed. The various embodiments of the restraining device 10 described herein may be available for temporary or chronic placement within a patient's body, and the restoration procedures performed through the use of an exemplary device 10 are reversible through minimally invasive procedures.

As shown in FIGS. 1 and 2, at least one embodiment of the restraining device 10 comprises a first engaging component 12 and a second engaging component 16. The first engaging component 12 may comprise a first shape and the second engaging component 16 may comprise a second shape that corresponds to at least a portion of the first shape of the first engaging component 12. For example, and without limitation, the first and second engaging components 12, 16 may be configured in a bar configuration as shown in FIGS. 1 and 2. Alternatively, the first and second engaging components 12, 16 may be configured in a contoured, circular or any other configuration suitable for use as referenced herein. Additionally, the first and second engaging components 12, 16 may be configured such that the shape of each of the components 12, 16 defines an interior area (not shown). It will be understood that the first and second engaging components 12, 16 of an exemplary restraining device 10 may be configured in any shape and may also be flexible, semi-flexible, or articulated. Furthermore, it is contemplated that a clinician may select the desired configuration of the components 12, 16 of the restraining device 10 based on the particular patient to be treated and/or pursuant to the specific application for which the restraining device 10 will be used to ensure that the restraining device 10 appropriately conforms to the tissue or

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organ of interest. For example, an exemplary embodiment of a restraining device 10 of the present disclosure may comprise first and second engaging components 12, 16 having a width of approximately 14 mm and a length of approximately 120 mm. In at least one additional embodiment, the first and second engaging components 12, 16 may have a width between 5 mm and 15 mm, and may have a length between 30 mm and 200 mm, with any number of potential widths and lengths contemplated by the present disclosure suitable for use with an exemplary restraining device 10 of the present disclosure.

In at least one embodiment of a restraining device 10 of the present disclosure, and as shown in FIG. 2, the first engaging component 12 of the restraining device 10 comprises a proximal end 13, a body having a first side 12A and a second side 12B, and a distal end 14. The first side 12A of the first engaging component 12 is configured to be positioned adjacent to or in contact with a tissue or organ of interest. Likewise, the second engaging component 16 comprises a proximal end 17, a body having a first side 16A and a second side 16B, and a distal end 18. The first side 16A of the second engaging component 16 is configured to be positioned adjacent to or in contact with the tissue or organ of interest.

The first engaging component 12 and the second engaging component 16 each comprise a material suitable to resist corrosion, such as and without limitation polyurethane, polyethylene, polytetrafluoroethylene ("PTFE"), nitinol, silastic, titanium, stainless steel or any other material suitable for use in the medical arts that is corrosion resistant. Accordingly, the restraining device 10 can withstand chronic placement within a body without the risk of deterioration. In at least one embodiment, the first and second engaging components 12, 16 of the restraining device 10 are comprised of ultra high density polyethylene.

The first and second engaging components 12, 16 of the restraining device 10 may be coupled to one or more springs 22, wherein the one or more springs 22 engage the first and second engaging components 12, 16 at their distal ends 14, 18 and proximal ends 13, 17, Springs 22, as well as struts 700 (as shown in FIGS. 7A and 7B) and couplers 1200 (as shown in FIGS. 14A through 16B), as referenced herein, may be generally referred to as "connector(s)" as they connect, in various embodiments, first engaging components 12 and second engaging components 16 of various restraining devices 10 of the present disclosure. The springs 22 may comprise torsion springs or any other type of spring known or developed that is capable of supporting the first and second engaging components 12, 16 a prescribed distance apart from each other, which may vary depending upon the orientation of the springs 22 about the first and second engaging components 12, 16. As described in more detail herein, due to the configuration and resistance of the springs 22, the first and second engaging components 12, 16 of the restraining device may be held such that little or no space is defined between the first sides 12A, 16A of the first and second engaging components 12, 16 based upon a first orientation of springs 22, and the first and second engaging components 12, 16 of the restraining device 10 may be held such that an interior space 70 is defined between the first sides 12A, 16A of the first and second engaging components 12, 16 based upon a second orientation of springs 22.

Further, each of the one or more springs 22 may have any number of coils configured in any fashion, provided the spring configuration and stiffness are suitable for the desired application of the restraining device 10. For example, and without limitation, in at least one embodiment each of the one or more springs 22 comprises a torsion, resistance spring

having three (3) closely-coiled coils. It will be appreciated that, in practice, a clinician may alter the number of coils and/or configuration of each spring 22 to achieve to a desired flexibility or rigidity of the springs 22 and, in this manner, the restraining device 10 may be customized for a particular patient and/or application for which the restraining device 10 is to be applied.

Each of the springs 22 of the restraining device 10 may comprise at least one rod 24 extending from each of the ends of its coils. The length of each rod 24 may be selected depending on the particular application for which the restraining device 10 is to be applied. As shown in the exemplary embodiments of a restraining device 10 in FIGS. 1 and 2, each rod 24 is coupled with either the first or second engaging component 12, 16. For example, as shown in FIG. 2, the first rod 24 of one of the springs 22 may be coupled with the second side 12B of the first engaging component 12 at or near a location on the distal end 14 of the first engaging component 12, and the second rod 24 of the same spring 22 may be coupled with the second side 16B of the second engaging component 16 at or near a location on the distal end 18 of the second engaging component 16. Furthermore, the rods 24 facilitate the formation of the interior space 70 (as shown in FIG. 2) between the first and second engaging components 12, 16 by holding the first and second engaging components 12, 16 in accordance with the tension of the springs 22.

The springs 22 of the restraining device 10 may comprise any dimensions so long as the restraining device 10 is of a sufficient size to move through a laparoscopic port and the springs 22 are capable of holding the first side 12A of the first engaging component 12 and the second side 16A of the second engaging component 16 a distance apart when the restraining device 10 has a configuration whereby the springs 22 are positioned about the first and second engaging components 12, 16. For example, springs 22 may have a “first” configuration as shown in FIG. 1, whereby springs 22 and the first and second engaging components 12, 16 are oriented in the same or substantially the same direction/plane so that the restraining device 10 may pass through a laparoscopic port. In such a first configuration, springs 22 engage the first and second engaging components 12, 16 so that the first and second engaging components 12, 16 either engage one another or have a relatively small space between them. Springs 22 may have a “second” configuration, for example and as shown in FIG. 2, when springs 22 are rotated about first and second engaging components 12, 16 and are not in the same direction/plane as the configuration of restraining device shown in FIG. 1. In such a second configuration, the first and second engaging components 12, 16 may define an interior space 70 when the restraining device 10 is “at rest,” with the interior space 70 increasing upon positioning the restraining device 10 around a stomach 100 (as shown in FIGS. 4A through 5A) for example, and potentially increasing only slightly further upon introduction of food, for example, into the stomach 100.

Accordingly, the dimensions of the springs 22 may dictate the native value of the interior space 70 between the first and second engaging components 12, 16. In at least one embodiment, the springs 22 may comprise a maximum outside diameter that is less than about 14 millimeters to allow a “collapsed” or “compressed” restraining device 10 to pass within a 15 mm diameter abdominal port. Furthermore, each of the springs 22 may comprise any material having a strength that is consonant with the application for which the restraining device 10 will be employed. In at least one embodiment, the springs 22 are comprised of a rigid or semi-rigid material that is suitable to resist corrosion, such as and without limitation,

polyurethane, PTFE, nitinol, silastic, titanium, stainless steel or any other material suitable for use in the medical arts that is corrosion resistant.

Referring back to FIG. 2, and as previously described, the springs 22 of the restraining device 10 are biased to maintain the first and second engaging components 12, 16 at a prescribed distance apart under a native, and as described above, “second” configuration. In this manner, when no pressure is applied to an exemplary restraining device 10—or when the restraining device 10 is “at rest”—the interior space 70 having a prescribed value is formed between the first and second engaging components 12, 16. The value of the interior space 70 may, in at least one embodiment, correlate with the outside diameter of the springs 22, minus the thickness of the first and second engaging components 12, 16. It will be understood that the value of the interior space 70 can be manipulated by a clinician depending on the thickness of tissue and/or organ to be treated or other factors. For example, to achieve an interior space 70 having a larger prescribed value, the outer diameter of the spring 22 may be increased and/or the thickness of the first and/or second engaging components 12, 16 may be adjusted. Accordingly, a clinician can easily modify the restraining device 10 such that it may be optimally configured for a particular application on a particular tissue.

Due to the nature of the springs 22 (depending on the material and rigidity selected), the first and second engaging components 12, 16 may exhibit some degree of “give” such that the first and second engaging components 12, 16 are capable of moving relative to each other when pressure is applied. While the first and second engaging components 12, 16 are biased to return to their resting position such that the interior space 70 is substantially equivalent to its prescribed/native value when no pressure is applied to the springs 22, the springs 22 of the restraining device 10 are also capable of allowing the first and second engaging components 12, 16 to move in response to force applied thereto. For example, and without limitation, if a force is applied against the first sides 12A, 16A of the first and second engaging components 12, 16, this force is translated to the springs 22 which enables the components 12, 16 of the restraining device 10 to move apart such that the interior space 70 is increased. Furthermore, if a force is applied directly to the springs 22 such that the coils are twisted about their axis in a direction counter to the coil configuration, similar to the above-listed example, the resultant effect on the restraining device 10 is that the first and second engaging components 12, 16 are moved apart and the interior space 70 is increased. However, due to the inherent bias provided by the springs 22 of the restraining device 10, after the pressure affecting the springs 22 is released, the springs 22—and thus the components 12, 16—return to their resting positions such that the interior space 70 reverts to its original prescribed value. It will be appreciated that the specific configuration and/or materials comprising each of the springs 22 may be selected to achieve the desired degree of elasticity depending on the application for which the restraining device 10 is to be applied.

As shown in FIG. 2, each rod 24 of spring 22 is rotatably coupled with the relevant component 12, 16 of the restraining device 10 such that each of the rods 24 is capable of pivotal movement with respect to the relevant component 12, 16 of the restraining device 10. The rotational coupling of the rods 24 with the first and second engaging components 12, 16 may be achieved through any means known in the art. For example and without limitation, the end of each rod 24 may comprise a pin that is capable of insertion into and rotation within a hole formed within the respective second side 12B, 16B of the applicable component 12, 16.

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As each of the rods **24** is coupled with a spring **22**, when the rods **24** rotate with respect to the components **12**, **16**, this effectively enables the respective spring **22** to move in an orbital manner with respect to the proximal or distal end of the restraining device **10**. Accordingly, the springs **22** are capable of rotating between a position that is substantially parallel with the first and second engaging components **12**, **16** (as shown in FIG. 1, referred to as a “first” configuration herein) and a position that is substantially perpendicular to the first and second engaging components **12**, **16** (as shown in FIG. 2, referred to as a “second” configuration).

The position of the springs **22** relative to the first and second engaging components **12**, **16** has the potential to significantly affect the overall width of the restraining device **10**. For example and without limitation, when the springs **22** are positioned in the substantially perpendicular position (i.e., a “second” configuration), the restraining device **10** may have a width that is more than twice the width of the same restraining device **10** when its springs **22** are positioned in the substantially parallel position. Accordingly, the rotational coupling of the rods **24** with the first and second engaging components **12**, **16** provides a clinician with the ability to manipulate the overall width of the restraining device **10** during laparoscopic delivery and/or implantation and further enables the restraining device **10** to be applied to tissues and/or organs having a length that is longer than the length of the restraining device **10** (as the proximal ends **13**, **17** and the distal ends **14**, **18** of the first and second engaging components **12**, **16** of the restraining device **10** are not obstructed by the springs **22**). In at least one embodiment, the widest part of the restraining device **10** is less than about 15 millimeters when the springs **22** are in the substantially parallel position such that the restraining device **10** can be easily inserted into a body cavity through a 15 millimeter trocar or port.

Now referring to FIG. 3A, a top view of at least one embodiment of a restraining device **10** of the present disclosure is shown. Here, the first and/or second engaging components **12**, **16** of the restraining device **10** further comprise one or more pads **30** extending therefrom. As shown in FIG. 3A, the one or more pads **30** may extend in a perpendicular fashion from the second side(s) **12B**, **16B** of the components **12**, **16** and/or in a lateral fashion from the components **12**, **16**. Each of the pads **30** are comprised of a flexible material and configured to provide an anchor through which sutures or any other type of anchoring device may be inserted. For example, and in at least one embodiment, one or more pads **30** may be comprised of polyurethane or any other material that is capable of securely holding sutures or another type of anchoring device therein. One or more sutures may be placed within suture apertures **32** as shown in FIG. 3A, whereby said suture apertures **32** may be positioned along pads **30** and/or one or both of components **12**, **16** as shown in FIG. 3A. Accordingly, after the first and second engaging components **12**, **16** are properly positioned about a targeted tissue or organ, sutures may be inserted through suture apertures **32** and secured to a superficial layer of the underlying targeted tissue or organ to assist in anchoring the restraining device **10** in the proper location thereon.

FIGS. 3B-3E show various embodiments of first engaging component **12** and/or second engaging component **16** with various features to facilitate suturing and/or connection of first engaging component **12** to second engaging component **16**. FIG. 3B shows an exemplary embodiment if a first and/or second engaging component **12**, **16** comprising suture apertures **32** positioned therethrough. Suture apertures **32** are not limited to positioning near one or more of the ends of said first and/or second engaging component **12**, **16**, as such suture

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apertures **32** may be positioned along first and/or second engaging component **12**, **16** as desired.

FIG. 3C shows an exemplary embodiment of a first and/or second engaging component **12**, **16** comprising one or more strings **34** affixed thereto. In the exemplary embodiment shown in FIG. 3C, two strings **34** are shown at or near the ends of first and/or second engaging component **12**, **16**, noting, however, that one, two, or more strings **34** may be affixed thereto and positioned as desired. In an exemplary embodiment of a restraining device **10**, and as shown in FIG. 4B, the first engaging component **12** and the second engaging component **16** may each comprise at least one string **34**, whereby said strings **34** may be connected to one another to facilitate a desired placement of restraining device **10** about a stomach **100**.

Additional embodiments of exemplary first and/or second engaging components **12**, **16** are shown in FIGS. 3D and 3E. As shown in FIG. 3D, first and/or second engaging component **12**, **16** comprises one or more suture members **36** affixed thereto, with said suture members defining an aperture to which a suture (not shown) or other restraining component may be affixed thereto. FIG. 3E shows an exemplary first and/or second engaging component **12**, **16** of the present disclosure comprising pads **30** positioned at or near each end of the first and/or second engaging component **12**, **16** with each pad **30** defining a suture aperture **32** therethrough. Additional embodiments of first and/or second engaging component **12**, **16** comprising features to allow for a suture or other restraining component to be positioned therethrough are also contemplated by the present disclosure. For purposes of the present disclosure, the term “suturing elements” shall mean any number of elements for introducing a suture into the first and/or second engaging component **12**, **16** and/or one or more pads **30** affixed thereto, including, but not limited to, one or more apertures **32**, strings, **34**, and/or suture members **36**. In addition, any number of the features of the various exemplary first and/or second engaging components **12**, **16** shown in FIGS. 3A-3E may appear in any number of embodiments of restraining devices **10** of the present disclosure.

In operation, the restraining device **10** may be applied to an organ or tissue of interest in order to restore the underlying tissue or organ into a desired configuration and/or provide support to the same. As discussed in further detail below, the restraining device **10** may be used for temporary or chronic implantation within a body without the risk of the first and second engaging components **12**, **16** migrating through or damaging the underlying tissue. Furthermore, because the restraining device **10** does not require that the underlying tissue be punctured in any significant manner to achieve restoration and/or provide support thereto, implantation of the restraining device **10** is entirely reversible and, if desired, the restraining device **10** may be easily removed from the organ or tissue of interest through a laparoscopic procedure.

As previously described, the specifications of the restraining device **10** may be modified to achieve a desired result. For example, and without limitation, the dimensions of the components **12**, **16** and/or the springs **22** may be chosen for a particular application and/or based on the patient. Accordingly, while certain embodiments of the restraining device **10** may be described in connection with particular tissues or organs, it will be appreciated that any of the embodiments of the restraining device **10** described herein may also be applied to any tissue or organ of interest in a similar manner and use of the particular embodiments of the restraining device **10** in lieu of others may be determined based on the patient’s specifications, the specific application, and/or the tissue or organ in question.

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In practice, an exemplary restraining device **10** is capable of restoring and/or supporting an underlying tissue while avoiding constriction and the excessive compression thereof. For example, the restraining device **10** may be applied to a stomach **100** as shown in FIGS. **4A** and **4B**. In at least one embodiment, and as shown in FIG. **4A**, the springs **22** of the restraining device **10** are rotated to the substantially perpendicular position such that the interior space **70** is increased between the first and second engaging components **12**, **16**. Thereafter, the restraining device **10** may be advanced over the stomach **100** such that, for example, the first side **12A** of the first engaging component **12** is positioned adjacent to the anterior wall of the stomach **100** and the first side **16A** of the second bar **16** is positioned adjacent to the posterior wall of the stomach **100**. While the restraining device **10** is shown in FIG. **4A** in a longitudinal placement with respect to the stomach **100**, it will be understood that the restraining device **10** may alternatively be positioned in a horizontal configuration or an angular configuration with respect to the stomach **100**.

After the first and second engaging components **12**, **16** are positioned in the desired location with respect to the stomach **100**, the overall interior space **70** may either comprise its original native interior space **70** or a relatively larger interior space **70** due to the positioning of the restraining device **10** about the stomach **100**. Therefore, various embodiments of restraining device **10** do not operate to “clamp” the stomach **100** or any other tissue or organ, as restraining device **10** merely operates, when positioned around a stomach **100**, to provide a limited pressure, if any, to maintain a native size/shape of at least a portion of the stomach **100**. For example, and depending upon the prescribed value of the interior space **70** as desired by a clinician, the configuration of the springs **22** can be modified to achieve a restraining device **10** that either does not compress, or only loosely compresses, the sandwiched tissue between the first and second engaging components **12**, **16**. In a preferred embodiment, restraining device **10**, when positioned about a stomach **100**, does not provide any meaningful compressive pressure upon the stomach **100**, and may be held in place, for example, using one or more sutures as referenced herein. In this manner, the restraining device **10** can be employed to reversibly restore an organ or tissue without forming adhesions thereon and/or permanently restoring the same.

In at least one embodiment, and depending on the configuration of the springs **22**, the springs **22** may exhibit enough elasticity to enable the first and second engaging components **12**, **16** to move to some degree in conjunction with any movement of a tissue or organ positioned between the first and second engaging components **12**, **16** of the restraining device **10**. In this manner, the springs **22** can allow, for example, the restraining device **10** to accommodate any inherent movement in the stomach **100** such that application of the restraining device **10** does not completely inhibit the normal digestive function of the same. For example, the introduction of food into the stomach **100** may cause the outer dimensions of the stomach **100** to expand, whereby stomach **100** exerts a pressure upon restraining device **100**. Furthermore, the restraining device's **10** ability to accommodate any inherent movement in the underlying organ and/or tissue increases the likelihood that the restraining device **10** will remain in its desired location on the tissue and/or organ without sheering off or sliding therefrom.

In the event it is desired that the restraining device **10** is further secured to the underlying tissue, and as previously referenced herein, a clinician can employ sutures to assist with the secure implantation of the restraining device **10** in the desired location. In the at least one embodiment of the

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restraining device **10** comprising the one or more pads **30**, a clinician can secure the one or more pads **30** of the first and second engaging components **12**, **16** to the underlying tissue through the use of superficial sutures. In this manner, the superficial sutures can be affixed through the one or more pads **30** and the surface of the underlying tissue such that the one or more pads **30** assist with anchoring the restraining device **10** in position on the tissue of interest. Conversely, and in at least one embodiment, sutures may be introduced directly to the first and/or second engaging components **12**, **16** to secure a restraining device to a tissue or organ of interest as shown in FIGS. **3A-3E**.

FIG. **4B** shows a cross-sectional view of the stomach **100** of FIG. **4A** taken along A-A. As shown in FIG. **4B**, when the first and second engaging components **12**, **16** of the restraining device **10** are longitudinally positioned on the stomach **100**, the stomach **100** is “divided” into two portions—a first stomach portion **110** and a second stomach portion **112** as shown in FIGS. **4A** and **4B**. In the embodiment shown in FIGS. **4A** and **4B**, the first stomach portion **110** is relatively smaller than the second stomach portion **112**. However, and as shown in FIG. **5A**, the first stomach portion **110** and the second stomach portion **112** may be approximately the same size.

As the first stomach portion **110** receives ingested matter directly from the gastroesophageal junction **99** as shown in FIG. **4A**, the placement of the restraining device **10** as shown in FIG. **4B** thus inhibits the majority of ingested matter from moving into the second stomach portion **112**. Instead, and in at least one embodiment, such ingested matter is directed through the smaller, first stomach portion **110** and into the pyloric canal **101** (as shown in FIG. **4A**) where a significant portion of the ingested matter is evacuated from the stomach **100**. Due to the size of the first stomach portion **110**, the amount of food that the patient can consume at one time is significantly reduced and satiety is more quickly achieved.

While the delineation formed by the restraining device **10** between the first stomach portion **110** and the second stomach portion **112** is not leak-proof, the interior space **70** created between the first and second engaging components **12**, **16** comprises an area that is less than the diameter of a fully extended stomach **100** (as shown in FIG. **4B**). Accordingly, when the restraining device **10** is applied in a longitudinal fashion to a stomach **100**, restraining device **10** provides support to the anterior and posterior walls of the stomach **100**, with the expansion of stomach **100** exerting a force onto the first and second engaging components **12**, **16** of restraining device **10**, this preventing stomach **100** distension in that area. In this manner, most of the food matter received into the first stomach portion **110** through the gastroesophageal junction **99** is maintained therein and the patient exhibits the sensation of satiety earlier.

Referring back to FIG. **4A**, and in at least one exemplary embodiment, because the restraining device **10** does not extend along the entire length of the stomach **100**, an outflow tract **114** is formed caudally of the restraining device **10**. This outflow tract **114** allows the portion of ingested matter that flows from the gastroesophageal junction **99** into the second stomach portion **112** to be evacuated from the stomach **100** in a controlled manner and to proceed through normal digestion. In addition, the outflow tract **114** allows any food matter or enzymes residing within the second stomach portion **112** to evacuate the stomach **100**.

Additional embodiments of restraining devices **10** of the disclosure of the present application positioned about a stomach **100** are shown in FIGS. **5B** and **5C**. In the exemplary embodiment shown in FIG. **5B**, restraining device **10** com-

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prises first and second engaging components **12**, **16** which are “longer” than those shown in FIG. 5A, whereby distal end **14** of first engaging component **12** and distal end **18** of second engaging component **16** extend past stomach **100**. In this exemplary embodiment, strings **34** affixed to first and second engaging components **12**, **16** may be connected/tied to each other after restraining device **10** has been positioned about a stomach, maintaining restraining device **10** in place.

FIG. 5C shows restraining device **10** positioned about stomach **100** viewed in the direction of the arrow shown in FIG. 5B. As shown in FIG. 5B, the tying/coupling together of strings **34** at the distal ends **14**, **18** of first and second engaging components **12**, **16** may cause portions of the first and second engaging components **12**, **16** to angle towards one another as shown in FIG. 5C. In this particular embodiment, first and second engaging components **12**, **16** are flexible or semi-flexible, and the positioning of restraining device **10** about stomach **100** serves to isolate the first stomach portion **110** from fundus **90** and greater curvature **92**. This “funnel” effect, in addition to the embodiment of a restraining device **10** shown in FIG. 5D below, may also be accomplished by having springs **22** with relatively shorter rods **24** (or relatively smaller struts **700** as referenced in various embodiments herein) at one end of restraining device **10**, and by having springs **22** with relatively longer rods **24** (or relatively larger struts **700** as referenced in various embodiments herein) at the other end of restraining device **10**. In addition as to being substantially rigid as shown in FIGS. 1 and 2, and as shown in FIG. 5C, first and second engaging components **12**, **16** may have substantially rigid center portions and one or more flexible or semi-flexible ends, or first and second engaging components **12**, **16** may be flexible or semi-flexible along the entirety of said components **12**, **16**.

An additional embodiment of a restraining device **10** of the present disclosure positioned about stomach **100** is shown in FIG. 5D. As shown in FIG. 5D, and as also discussed herein regarding FIG. 10, restraining device **10** further comprises a pliable junction **1000** positioned within the first engaging component **12** and second engaging component **16**, effectively “splitting” each component, whereby each “split” component is connected to one another by way of a flexible structure **1002**. As shown in FIG. 5D, flexible structure **1002** may comprise a PTFE band or a PTFE mesh, for example, whereby any potential embodiment of flexible structure **1002**, using one or more pliable materials disclosed herein or otherwise known in the art, allows the relatively or fully rigid first engaging component **12** and second engaging component **16** to better fit the stomach, allowing the patient, for example, to bend the stomach when restraining device **10** is positioned thereon.

Approximating first and second engaging components **12**, **16**, as shown in FIGS. 5B, 5C, and 5D, and as identified during internal testing of exemplary restraining devices **10** of the present disclosure, leads to reduced food intake (65-70% of normal food intake) as compared to positioning restraining device **10** about stomach **100** without approximating first and second engaging components **12**, **16** toward one another as shown in FIG. 4A (80-85% of normal food intake). This approximation also assists with the prevention of a fistula so that the food may move down the pouch (first stomach portion **110**) along antrum **94** as opposed to shunted to the fundus **90** of stomach **100**.

As described herein, application of the restraining device **10** allows a clinician to restore a targeted tissue, such as a stomach **100**, while avoiding constriction and excessive compression of the same. Further, the various embodiments described herein allow a clinician to tailor the restraining

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device **10** to multiple restoration applications and various different types of tissues. Permanent restoration of the tissue is avoided, which prevents adhesions from developing in the underlying targeted tissue and allow for the complete reversal of the restoration procedure. Additionally, the restraining device **10** is simple to deliver and, as such, the device **10** may be used in conjunction with other techniques or surgical procedures.

Regarding the application of the restraining device **10** to the stomach **100**, use of the restraining device **10** in the treatment of obesity avoids the nutritional and metabolic deficiencies observed after Malabsorptive Procedures because the digestive process may continue within the stomach as with a native stomach **100**. In addition, the restraining device **10** does not require more than superficial punctures to the underlying tissue, nor does it employ pins, staples or significant sutures which may lead to dehiscence or fistula formation, or produce the degree of regurgitation and vomiting observed in connection with conventional methods used to treat obesity. Moreover, each of the embodiments described herein may be inserted into the body cavity laparoscopically, thereby decreasing the patient’s stress associated with the procedure and the patient’s recovery time. It will be recognized that any of the devices described herein may be employed in combination with other conventional bariatric procedures.

Now referring to FIG. 6, a flow chart of a method **300** for laparoscopically delivering the restraining device **10** is shown. For ease of understanding, the steps of the related methods described herein will be discussed relative to the components of an exemplary restraining device **10**. Furthermore, while the methods described herein are described in connection with embodiments of the restraining device **10** and an exemplary delivery device, it will be appreciated that various additional devices may be used to facilitate the laparoscopic delivery of the restraining device **10** such as a camera, light and/or a device for delivering a gas to a targeted area.

At step **302**, the first and second engaging components **12**, **16** of restraining device **10** are advanced laparoscopically into the patient’s body. In at least one embodiment, the restraining device **10** may be inserted through a 15 millimeter cannula under insufflation into the appropriate cavity of the patient’s body. This may be achieved through use of an exemplary delivery device known in the art. At this step **302**, the springs **22** of the restraining device **10** are positioned in the substantially parallel position such that the overall diameter of the restraining device **10** is sufficiently narrow for insertion into the body.

At step **304** the restraining device **10** is advanced to a location adjacent to a targeted tissue. In the at least one embodiment of the method **300** where an exemplary delivery device is employed to facilitate delivery of the restraining device **10** to the targeted tissue, at step **304** the restraining device **10** is advanced out of the delivery device and into the body cavity. After the restraining device **10** is no longer positioned within an exemplary delivery device, the delivery device may be withdrawn from the body cavity at this step **304** or as desired by the clinician.

At step **306**, the springs **22** of the restraining device **10** are rotated from the substantially parallel position to the substantially perpendicular position, separating the first and second engaging components **12**, **16** from one another to a native interior space **70**. Step **306** may be performed using any number of standard laparoscopic tools known in the art useful to pull and grasp portions of a tissue or a device. In this manner, neither the proximal ends **13**, **17** nor distal ends **14**,

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18 of the first and second engaging components 12, 16 of the restraining device 10 are blocked by the springs 22 and/or rods 24, and the first and second engaging components 12, 16 may be advanced over a targeted tissue having a length that is greater than the overall length of the first and second engaging components 12, 16.

At step 308, under fluoroscopic, direct camera control or otherwise, the restraining device 10 is advanced over the targeted tissue. In at least one embodiment, and at step 308, the first side 12A of the first engaging component 12 is positioned adjacent to the desired surface of the targeted tissue and the first side 16A of the second engaging component 16 is positioned adjacent to an opposite side of the targeted tissue. As the first and second engaging components 12, 16 of the restraining device 10 are positioned adjacent to opposite sides of the targeted tissue, at this step 308 the targeted tissue is positioned within the interior space 70 formed between the first and second engaging components 12, 16. Accordingly, while the targeted tissue may experience some compressional force exerted by the first and second engaging components 12, 16 of the restraining device 10, the majority of the pressure upon the first and second engaging components 12, 16 is provided by distension/expansion of the targeted tissue (for example, expansion of a stomach 100 when food is introduced therein). Further, due to the configuration and composition of the restraining device 10, the restraining device 10 can remain within the patient's body for as long as the restoration or support treatment delivered thereby is desired.

If preferred, in at least one embodiment of the restraining device 10 that further comprises one or more pads 30 coupled with the first and/or second engaging components 12, 16, the method 300 may advance from step 308 to step 310. At step 310, a clinician can employ sutures to further anchor and secure the restraining device 10 in the desired position on the targeted tissue. These sutures need only superficially puncture the underlying tissue and therefore are not associated with the negative effects associated with suturing, stapling and/or the insertion of pins used in conventional methods.

Additional embodiments of restraining devices 10 of the disclosure of the present application is shown in FIGS. 7A and 7B. As shown in FIGS. 7A and 7B, restraining device 10 comprises a first engaging component 12 and a second engaging component 16, whereby the first engaging component 12 and second engaging component 16 are coupled to one another by way of one or more struts 700. In the exemplary embodiment shown in FIGS. 7A and 7B, four struts 700 and five struts 700 are used, respectively, but any number of struts 700 may be used as desired for a particular application. Struts 700 may comprise any number of suitable materials as otherwise described herein, including, but not limited to, nitinol and stainless steel. Struts 700, in an exemplary application, would have a pre-established "open" size and a pre-established "strength", so that restraining device, in a native "rest" configuration, would be "open" (by way of, for example, the "memory" of struts 700) and would require some sort of pressure/force to open even further. The "open" configuration would keep first engaging component 12 and second engaging component 16 a fixed distance apart to avoid gastric tissue compression when positioned about a stomach.

An exemplary restraining device 10 may further comprise a mesh curtain 702 coupled to struts 700 as shown in FIGS. 7A and 7B. Mesh curtain 702 may prevent or limit potential organ distension or remodeling when restraining device 10 is positioned about an organ of interest. A demi shaft 704, as shown in FIGS. 7A and 7B, may be positioned around at least part of struts 700 and/or mesh curtain 702 to facilitate insertion of restraining device 10 within a body. For example, and

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as shown in FIG. 7A, restraining device 10 may be seen as inserted within a body, noting that restraining device is somewhat compressed by demi shaft 704, to facilitate insertion through, for example, a laparoscopic port. After insertion, and as shown in FIG. 7B, demi shaft 704 may be withdrawn in the direction of the arrow (shown as partially withdrawn in the figure) to allow restraining device 10 to obtain its original, uncompressed configuration, and ultimately removed from the body. Struts 702 may have a first configuration when positioned within demi shaft 704, as shown in FIG. 7A, and may have a second configuration when demi shaft 704 is removed, as shown on the left side of FIG. 7B.

When restraining device 10 is in its native configuration, it may then be positioned about a stomach, for example, and the user positioning restraining device 10 may grasp stomach tissue using any number of laparoscopic tools through mesh aperture 706 as shown in FIGS. 7B and 8. As shown in FIG. 8, an exemplary restraining device 10 may also comprise one or more suture members 36 to allow one or more sutures (not shown) to be used to secure restraining device 10 to, for example, an anterior gastric wall.

An additional embodiment of a restraining device 10 of the disclosure of the present application is shown in FIG. 9. As shown in FIG. 9, restraining device 10 comprises a first engaging component 12 and a second engaging component 16, whereby first engaging component 12 and second engaging component 16 are coupled to one another by way of one or more springs 22. In this exemplary embodiment, restraining device 10 further comprises a mesh curtain 702 coupled to first engaging component 12 and second engaging component 16, whereby mesh curtain 702 may prevent or reduce organ distension or remodeling when restraining device 10 is positioned about an organ. The flexibility/pliability of mesh curtain 702 would allow mesh curtain 702 to be closely positioned first engaging component 12 and second engaging component 16 upon insertion of restraining device 10 within a body, and may further allow mesh curtain 702 to expand in a direction of springs 22 as shown in FIG. 9, for example, to prevent organ distention.

FIG. 10 shows yet another embodiment of a restraining device 10 of the present application. As shown in FIG. 10, restraining device comprises a first engaging component 12 and a second engaging component 16, whereby first engaging component 12 and second engaging component 16 are coupled to one another by way of one or more springs 22. However, in this exemplary embodiment, restraining device 10 further comprises a pliable junction 1000 positioned within the first engaging component 12 and second engaging component 16, effectively "splitting" each component, whereby each "split" component is connected to one another by way of a flexible structure 1002. As shown in FIG. 10, flexible structure 1002 may comprise a PTFE band or a PTFE mesh, for example, whereby any potential embodiment of flexible structure 1002, using one or more pliable materials disclosed herein or otherwise known in the art, allows the relatively or fully rigid first engaging component 12 and second engaging component 16 to better fit the stomach, allowing the patient, for example, to bend the stomach when restraining device 10 is positioned thereon. In addition, and as shown in the exemplary embodiment in FIG. 10, restraining device 10 may further comprise a tissue cover 1004 positioned either partially or fully around either or both of first engaging component 12 and second engaging component 16.

FIG. 11 shows an exemplary embodiment of a restraining device 10 of the present application positioned within a laparoscopic port. As shown in FIG. 11, restraining device 10 is shown as being inserted into a body portion by way of lap-

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aroscopic port 1100, with the exemplary laparoscopic port 1100 shown as comprising a ring 1102 and a port sleeve 1104 to facilitate introduction of restraining device 10 through an abdominal wall 1106. Viewing the figure from the outside-in, restraining device 10 is shown as being introduced into an abdomen through abdominal wall 1106, with peritoneum 1108 shown as being an innermost layer of the abdominal wall.

An additional embodiment of a restraining device 10 of the disclosure of the present application is shown in FIG. 12A. As shown in FIG. 12A, restraining device 10 comprises a first engaging component 12 and a second engaging component 16, whereby first engaging component 12 and second engaging component 16 are coupled to one another by way of one or more couplers 1200. In this exemplary embodiment, restraining device comprises two couplers 1200, shown in further detail in FIG. 12B. Couplers 1200 may comprise coupler arms 1202 connected to one another by way of a coupler bar 1204, and may comprise any number of suitable materials as described herein for various components of the present disclosure. As shown in FIG. 12A, coupler arms 1202 may be positioned within first engaging component 12 and second engaging component 16 by way of apertures 1210 (as shown in FIG. 12B) defined therein, allowing first engaging component 12 and second engaging component 16 to move relative to one another as couplers 1200 swivel when positioned at least partially within said apertures 1210.

As shown in FIG. 12B, an exemplary embodiment of a coupler 1200 may comprise a coupler protrusion 1206 at or near the end of each coupler arm 1202, whereby coupler protrusions either prevent or restrict the removal of coupler 1200 from the either first engaging component 12 or the second engaging component 16 by way of stops 1208 positioned about the first engaging component 12 and the second engaging component 16 at or near the apertures 1210 defined therein. As coupler 1200 moves in the direction of the arrow shown in FIG. 12B, coupler protrusions 1206 would engage stops 1208, thus preventing or restricting the removal of coupler 1200 from the first engaging component 12 and/or the second engaging component 16.

Insertion of the exemplary embodiment of the restraining device 10 shown in FIG. 12A into a body portion may be performed by inserting said restraining device 10 through a laparoscopic port with first engaging component 12 and second engaging component 16 either touching or nearly touching one another, and with couplers 1200 either mostly or fully inserted within apertures 1210 of restraining device 10. When restraining device has been inserted into a body portion, first engaging component 12 and second engaging component 16 may be separated a pre-established distance from one another by way of turning/swiveling couplers 1200. As such, couplers 1200, as well as first engaging component 12 and second engaging component 16, may be sized and shaped as desired for a particular application.

FIGS. 13A and 13B show an exemplary embodiment of a restraining device 10 with a coupler 1200 positioned therein in a configuration to maximize the interior space 70 between first engaging component 12 and second engaging component 16. As such, the configuration of coupler 1200 shown in FIG. 12A may be viewed as “closed,” while the configuration shown in FIG. 13A may be viewed as “open.” In addition, and as described above and shown in FIG. 13A, coupler 1200 may be withdrawn from apertures of first engaging component 12 and second engaging component 16 to facilitate placement of restraining device about an organ of interest. Such withdrawal may be performed using any number of laparoscopic tools,

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for example, whereby coupler 1200 is withdrawn up to a point where protrusions 1206 would engage stops 1208 as shown in FIG. 12B.

Another exemplary embodiment of a restraining device 10 of the disclosure of the present application is shown in FIGS. 14A and 14B. As shown in FIGS. 14A and 14B, restraining device 10 comprises a first engaging component 12 and a second engaging component 16, whereby first engaging component 12 and second engaging component 16 are coupled to one another by way of one or more couplers 1200. In the embodiments shown in FIGS. 14A and 14B, couplers 1200 comprise “scissor” couplers 1200, whereby at least two coupler bars 1204 are connected to one another via a pivot member 1400 so that coupler bars 1204 may pivot about one another at pivot member 1400. As coupler 1200 shifts from a “closed” configuration, as shown in FIG. 14A, to an “open” configuration, as shown in FIG. 14B, coupler bars 1204 pivot about pivot member 1400 causing coupler bars 1204 to move closer to one another in a direction opposite the direction of the length of first engaging component 12 and second engaging component 16.

Couplers 1200, as shown in FIGS. 14A and 14B, may “open” and “close” by way of movement of coupler arms 1202 (ends shown in the two figures) within apertures 1210 defined within first engaging component 12 and second engaging component 16. Apertures 1210, as shown in these exemplary embodiments, are configured as horizontal grooves to allow coupler arms 1202 to move closer to one another within the same aperture 1210 while coupler 1200 is “opening” and to allow coupler arms 1202 to move away from one another within the same aperture 1210 while coupler 1200 is “closing.” As shown in FIG. 14B, restraining device 10, in a fully open position, would maximize the interior space 70 between first engaging component 12 and second engaging component 16. Protrusions 1402, such as teeth, indentations, and the like, may be positioned at or near apertures 1210, as shown in FIG. 14B, to facilitate fixation of coupler arms 1202 in a desired position, allowing a user positioning restraining device 10 within a body to measure the exact the interior space 70 between first engaging component 12 and second engaging component 16.

Upon insertion of the exemplary embodiment of restraining device 10 shown in FIGS. 14A and 14B within an abdominal cavity (through a laparoscopic port, for example), any number of laparoscopic tools may be used to grasp couplers 1200 to withdraw them from first engaging component 12 and second engaging component 16 as referenced herein regarding various other embodiments of restraining devices 10.

An exemplary delivery apparatus for delivering a restraining device 10 of the present application is shown in FIGS. 15A and 15B. As shown in FIG. 15A, an exemplary delivery apparatus 1500 comprises a shaft 1502 and arms 1504 at or near a distal end 1506 of shaft 1502. Apparatus 1500 may further comprise string 1508, whereby string 1508 may either be coupled to or engaged by a pull bar 1510 (an exemplary “puller” of the present disclosure) positioned within the lumen of apparatus 1500. Arms 1504 may engage an outer surface of first engaging component 12 and second engaging component 16, a protrusion (not shown) extending from said components, or by way of a groove 1512 (shown in FIG. 15B) positioned within said components.

As shown in FIG. 15A, string 1508 is coupled to springs 22 of restraining device 10, whereby a “pulling” motion of pull bar may effectively pull springs 22 from a first configuration (not shown in FIG. 15A) to a second configuration as shown in FIG. 15A so that restraining device may be positioned about an organ of interest. In an additional embodiment,

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string 1508 may be positioned at least partially within the lumen of apparatus 1500, whereby a user of apparatus may pull string 1508 instead of pulling pull bar 1510 to operate apparatus 1500. Various components of apparatus 1500 may comprise any number of suitable materials, including, but not limited to, the various materials referenced herein in connection with components of restraining device 10. For example, string 1508 may comprise, for example, plastic or metal thread.

For example, and in an abdominal cavity with the epiploon lesser curvature dissected, arms 1504 of apparatus 1500 may position restraining device 10 to a desired location, springs 22 of restraining device 10 may be widely opened, thus separating the first engaging component 12 and the second engaging component 16 by using pull bar 1510 to pull the strings 1508 and move springs 22 from an axial position to a 90° position relative to first engaging component 12 and the second engaging component. An “opened” restraining device 10 may then be introduced through the dissected lesser curvature space in parallel position to the intragastric mannequin tube, thus creating a desired pouch size. First engaging component 12 and second engaging component 16, upon positioning restraining device 10 about a stomach, would occupy the anterior and posterior wall of the stomach. When the first engaging component 12 and the second engaging component 16 are located in the desired position, spring 22 approximates the first engaging component 12 and the second engaging component 16 to each other gradually without producing any ischemic or tissue damage. Upon moving springs 22 to their desired location and/or positioning restraining device 10 about the stomach, restraining device 10 is separated from apparatus 1500, strings 1508 are cut and apparatus 1500 is pulled out of the abdominal cavity through the port. Restraining device 10 may then be secured in the gastric tissue with superficial stitches.

FIG. 15B shows an exemplary embodiment of at least a portion of apparatus 1500 engaging a “closed” restraining device 10. As shown in FIG. 15B, arms 1504 of apparatus 1500 are shown engaging first engaging component 12 and second engaging component 16 of restraining device 10 by way of grooves 1512 positioned within said components, permitting apparatus 1500 to deliver restraining device 10 through an abdominal port.

FIGS. 16A and 16B show an exemplary embodiment of an apparatus 1500 engaging/positioning an exemplary restraining device 10 of the disclosure of the present application. As shown in FIGS. 16A and 16B, arms 1504 of apparatus 1500 may engage the first engaging component 12 and the second engaging component 16 of restraining device 10, whereby the “opening” of arms 1504 would facilitate the “opening” of restraining device from a “closed” configuration (as shown in FIG. 16A) to an “open” configuration (as shown in FIG. 16B).

“Opening” restraining device 10, as shown in FIG. 16B, may be performed as follows. In the abdominal cavity with the epiploon lesser curvature dissected, the arms 1504 of apparatus 1500 may be widely opened, thus separating the first engaging component 12 and the second engaging component 16 of restraining device 10. A surgeon, for example and with the aid of laparoscopic graspers, may then “pull” out couplers 1200, placing couplers 1200 in a withdrawn configuration as shown in FIG. 16B. Restraining device 10, in an “open” configuration, may be introduced through the dissected lesser curvature space in parallel position to the intragastric mannequin tube, thus creating the desired stomach pouch size. First engaging component 12 and second engaging component 16, when positioned about a stomach, would occupy the anterior and posterior wall of the stomach. When the first engaging component 12 and the second engaging component 16 are

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located in the desired position, the user/operator may then approximate the plates to each other gradually (millimetrically) using apparatus 1500 device without producing any ischemic or tissue damage. When the user/operator has positioned restraining device 10, apparatus 1500 would then be separated from restraining device and pulled out of the abdominal cavity through the port. If desired, restraining device may also be secured in the gastric tissue with superficial stitches.

FIGS. 17A and 17B show another exemplary embodiment of an apparatus 1500 engaging/positioning an exemplary restraining device 10 of the disclosure of the present application. As shown in FIGS. 17A and 17B, arms 1504 of apparatus 1500 may engage the first engaging component 12 and the second engaging component 16 of restraining device 10, whereby the use of string 1508 connected to the first engaging component 12 and the second engaging component 16 may be used by an operator of apparatus 1500 to “open” and/or “close” said components. One or more strings 1508, in such exemplary embodiments, may be coupled at their first ends to the first engaging component 12 and/or the second engaging component 16, and may be coupled at their second ends to a string rotator 1700 (another exemplary “puller” of the present disclosure) as shown in FIGS. 17A and 17B. Operation/rotation of string rotator 1700 may then facilitate the “opening” of restraining device 10 from a “closed” configuration (as shown in FIG. 17A) to an “open” configuration (as shown in FIG. 17B). In such an embodiment, arms 1504 may also be positioned upon the first engaging component 12 and/or the second engaging component 16 at or near the center of said components (forming a “pivot” area), and the string rotator 1700 and strings 1504 affixed thereto may operate to keep said components in their axial position. Ultimate restraining device 10 delivery may be performed using one or more methods described herein.

Restraining device 10 may also be positioned “perpendicularly” using apparatus 1500 as shown in FIG. 18. As shown in FIG. 18, arms 1504 and strings 1508 are shown positioned about the first engaging component 12 and the second engaging component 16 “perpendicularly” so that restraining device 10 and apparatus 1500 are not aligned with one another. Rotation of string rotator 1700, as described above, may facilitate the opening and/or closing of restraining device 10.

The various embodiments of restraining devices 10 of the present disclosure may be introduced into a body cavity through, for example, a laparoscopic port. Such restraining devices 10 would be inserted while “compressed” (as shown in FIGS. 1, 7A, 10, 11, 12A, and 14A, for example), and when inserted, would be “opened” or “deployed” (as shown in FIGS. 2, 7B, 8, 9, 13A, 13B, and 14B, for example) prior to being positioned about a stomach 100. In addition, a user of such a restraining device 10 may adjust the opening/deployed dimensions of restraining device 10 prior to positioning it about a stomach, and may further tailor the open/deployed dimensions of restraining device 10 after it has been positioned as desired.

Furthermore, any number of exemplary embodiments of restraining devices 10 of the present application may be wholly or partially resorbable by the body. For example, and using the exemplary restraining device 10 shown in FIGS. 1 and 2 as an example, first and second engaging components 12, 16 may not be resorbable by the body, while springs 22 may be resorbable. In such an embodiment, restraining device 10, once positioned about a stomach 100, may slowly begin the process of total or partial resorption. Resorbable materials suitable for one or more portions of restraining devices 10

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may include, but are not limited to, polyglycolide (PGA), polylactide (PLA), L-lactide (LPLA), poly(DL-lactide) (DL-PLA), poly(ε-caprolactone) (PCL), poly(dioxanone) (PDO), polyglycolide-trimethylene carbonate (PGA-TMC), poly(D, L-lactide-co-glycolide) (DLPLG), or combinations thereof.

In a situation where it is desired to have restraining device 10 serve as a reversible bariatric device, and if it is desired not to engage in a subsequent procedure to, for example, laparoscopically remove some or all of restraining device 10 from a body, some or all of restraining device 10 may be resorbed. By way of example, an exemplary restraining device comprising resorbable springs 22 may be positioned about a stomach, and first and second engaging components 12, 16 may slowly become coated with various fibrotic tissue. As springs 22 resorb, springs 22 will eventually no longer operate to connect first and second engaging components 12, 16 to one another, which will effectively cause the various portions of restraining device 10 to no longer serve as a restraining mechanism. Springs 22 may also become coated, in part or in their entirety, by fibrotic tissue, so that when springs 22 resorb, first and second engaging components 12, 16 remain positioned about stomach 100, but the introduction of food into a stomach 100, for example, does not exert any pressure on springs 22 by way of first and second engaging components 12, 16 as first and second engaging components 12, 16 are no longer connected to one another by way of springs 22. Furthermore, first and second engaging components 12, 16 may be resorbable and springs 22 may not, so as first and second engaging components 12, 16 resorb within the body, springs 22 no longer serve to connect first and second engaging components 12, 16 to each other, and restraining device 10 no longer performs any restraining function. Additional embodiments of restraining devices 10 may be resorbable, including, but not limited to, struts 700 and mesh curtain 702.

Regarding removal of exemplary embodiments of restraining devices 10 of the present application, said restraining devices 10 may be removed in whole or in part, for example, by way of a laparoscopic procedure. Reversibility of restraining devices 10, as referenced in the present application, pertains to the ability to position a restraining device 10 about a tissue or organ, and at some time thereafter, have some or all of restraining device 10 resorb within a body or remove some or all of restraining device from the body so that restraining device 10 no longer functions to restrain a tissue or organ. For example, and referencing the exemplary embodiments shown in FIGS. 1 and 2, a laparoscopic procedure to remove springs 22 from the restraining device 10 and the body, but not to remove first and second engaging components 12, 16 from the body, would have the effect of “reversing” the procedure of placing said restraining device 10 about a tissue or organ. Similarly, and as shown in FIGS. 7A, 7B, and 8, removal of struts 700 from restraining device 10 would also have the effect of “reversing” the placement of restraining device 10 about a tissue or organ.

Furthermore, and if desired, the entirety of restraining device 10 may be removed laparoscopically by, for example, reversing the steps used to insert said restraining device 10 within a body. For example, a method 1900 for removing a restraining device 10 from a body as shown in FIG. 19 may comprise the steps of withdrawing restraining device 10 from a tissue or organ (withdrawal step 1904), configuring restraining device 10 to fit within a laparoscopic port (configuration step 1906), and removing restraining device 10 from the body through a laparoscopic port (removal step 1908). Such an exemplary method may be preceded by positioning a laparoscopic port within a body to facilitate removal of restraining device 10 (port insertion step 1902), and may be followed by

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removing said laparoscopic port from the body after removal step 1908 (port removal step 1910).

Additional embodiments of restraining devices 10 of the present disclosure are shown in FIGS. 20 and 23. As shown in the embodiments shown in FIGS. 20 and 23, restraining devices 10 comprise a first engaging component 12 and a second engaging component 16, such as those provided in various other embodiments herein. First engaging component 12 and second engaging component 16, in such an exemplary embodiment, are positioned relatively parallel to one another. A first swivel arm 2000 and a second swivel arm 2002 are pivotally connected to first engaging component 12 and second engaging component 16, so that first swivel arm 2000 engages first engaging component 12 at pivot point 2004 and engages second engaging component 16 at pivot point 2006, and so that second swivel arm 2002 engages first engaging component 12 at pivot point 2008 and engages second engaging component 16 at pivot point 2010. First swivel arm 2000 and second swivel arm 2002 may swivel in the directions shown in arrows A and B in FIGS. 20 and 23, noting that first swivel arm 2000 and second swivel arm 2002 may move in opposite directions, so that one moves in the direction of arrow A and the other moves in the direction of arrow B, or first swivel arm 2000 and second swivel arm 2002 may move in the same direction.

First engaging component 12 and/or second engaging component 16 may present studs 2012, as shown in FIGS. 20 and 23, whereby studs 2012 permit first swivel arm 2000 and/or second swivel arm 2002 to be engaged thereto and permit swiveling/rotation about the various pivot points. In at least one embodiment, studs 2012 have a bulbous component relatively larger than a non-bulbous portion, so that a first swivel arm 2000 and/or a second swivel arm 2002 presenting one or more apertures 2014 therethrough may engage studs 2012. Any number of other means to couple first swivel arm 2000 and/or second swivel arm 2002 to first engaging component 12 and/or second engaging component 16 may be used, such as pins, rivets, snaps, screws, and/or other fasteners or coupling means.

First swivel arm 2000 and second swivel arm 2002, as shown in FIGS. 20 and 23, may further comprise a first interconnection arm 2016 and a second interconnection arm 2018 positioned relative to bends 2020, 2022 of said arms. Bends 2020, 2022, as shown in FIGS. 20 and 23, are positioned in the relative middle (somewhere between the relative ends) of first swivel arm 2000 and a second swivel arm 2002, so that first swivel arm 2000 and second swivel arm 2002 form a relative “U-shape” within restraining device 10. First swivel arm 2000 and/or a second swivel arm 2002 may further comprise various additional curvatures 2024, such as shown in FIG. 20, such that bends 2020, 2022 and/or curvatures 2024 define a native U-shaped configuration as previously described.

An interior space 70 between the first engaging component 12 and the second engaging components 16, as shown in FIGS. 20 and 23, may change depending on various factors and/or configurations of restraining device 10. For example, and during insertion of an embodiment of a restraining device 10 such as shown in FIG. 20 or 23 through a laparoscopic port, first swivel arm 2000 and a second swivel arm 2002 may be relatively parallel to, or in the same relative plane as, first engaging component 12 and second engaging component 16, so that the interior space 70 is relatively small. After insertion, and after swiveling first swivel arm 2000 and a second swivel arm 2002 about first engaging component 12 and second engaging components 16, the interior space 70 may either stay the same or increase to facilitate placement of restraining

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device **10** about a stomach, for example. Furthermore, the use of a tape **2026** positioned about one or more of the first swivel arm **2000** and the second swivel arm **2002**, as shown in FIG. **20**, may further adjust the amount of interior space **70** between the first engaging component **12** and the second engaging components **16** depending on the amount of relative tension (pull) is applied to tape **2026**. For example, if tape **2026** is positioned about one or more of the first swivel arm **2000** and the second swivel arm **2002** so that pulling tape **2026** in a direction shown by arrow C in FIG. **20** causes portions of tape **2026** to press against first interconnection arm **2016** and/or second interconnection arm **2018** causing the relative ends of each of first swivel arm **2000** and a second swivel arm **2002** to move toward each other, the amount of interior space **70** between first engaging component **12** and second engaging component **16** may be reduced. In such an embodiment, restraining device **10** could be inserted into a mammalian body, positioned around a bodily organ without applying pressure to said organ, and tape **2026** could be pulled so that the amount of interior space **70** is adjusted as desired for the particular application. In a bariatric application, for example, it is desired not to have portions of restrictive device **10** apply any pressure about the stomach, but tape **2026** could be pulled/adjusted so first engaging component **12** and second engaging component **16** properly engage the stomach and be sutured thereto as desired. Distention of the stomach, such as by the introduction of solid and/or liquid food or water into the stomach, would cause the stomach to exert a force/pressure against the first engaging component **12** and second engaging component **16**, forming a first stomach portion **110** and a second stomach portion **112** as shown in FIGS. **4A** and **4B** referenced above and FIG. **21** referenced below.

In addition, and as shown in the exemplary embodiments of restraining devices **10** of the present disclosure shown in FIGS. **20** and **23**, various portions of restraining device **10**, such as the first engaging component **12**, the second engaging component **16**, the first swivel arm **2000**, and/or the second swivel arm **2002** may have one or more suture apertures **32** defined therethrough to facilitate the placement of one or more sutures **34** (also referred to herein as strings **34**) to couple restraining device **10** to a portion of a mammalian body, such as a stomach. In various embodiments, one or more of the relative ends of the first engaging component **12** and the second engaging component **16** may be rounded and/or tapered to improve overall patient comfort when a restraining device is positioned within the patient's body.

FIG. **21** shows an exemplary embodiment of a restraining device **10**, such as shown in FIG. **20**, positioned about a stomach. As shown in FIG. **21**, device **10** may be positioned about stomach **100** similar to placement of restraining device **10** about stomach **100** as shown in FIG. **5B**, so that the first engaging component **12** and the second engaging component **16** (not shown in FIG. **21**) may engage opposite sides of stomach **100**. Restraining device **10** may be secured to stomach **100** using one or more sutures **34** to couple restraining device **10** to stomach **100**, and restraining device **100** may be further positioned about stomach **100** by way of adjusting tape **2026** positioned around restraining device **10**. As shown in FIG. **21**, tape **2026** is positioned about restraining device **10** so to engage first interconnection arm **2016** and second interconnection arm **2018**, whereby pulling tape **2026** in a direction shown by arrow C would cause first engaging component **12** and second engaging component **16** to move closer to one another. Tape **2026** may have one or more detectable portions **2100** positioned/imprinted thereon, so that a user of restraining device **10** can "see" the adjustment of tape **2026** by way of fluoroscopy (if detectable portions **2100** are radiopaque),

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camera, or other means whereby portions of a device positioned within a body can be visualized, either directly or through some sort of technological means, to allow a user of such a device to adjust the same.

Furthermore, and as shown in FIG. **21**, restraining device **10** may have a cover flap **2012** positioned thereon to assist maintaining the placement of restraining device **10** about stomach **100**. Cover flap **2012**, as shown in FIG. **21**, may be ultimately coupled to first engaging component **12** and second engaging component **16** (not shown in FIG. **21**), so that after restraining device **10** is positioned about a stomach **100**, cover flap **2012** may be closed (initially secured to first engaging component **12** and closed by way of securing cover flap **2012** to second engaging component **16**, or vice versa) about an upper portion of stomach **100**.

As referenced above, restraining device **10** is positioned about stomach **100** so not to exert pressure upon stomach **100**, but when stomach **100** becomes distended, stomach **100** exerts pressure/force upon first engaging component **12** and second engaging component **16** of restraining device **10**.

Steps of a method for delivering and positioning an embodiment of a restraining device **10**, such as the embodiment shown in FIGS. **20** and **21**, are shown in FIG. **22**. As shown in FIG. **22**, exemplary method **2200** comprises the step of laparoscopically inserting/advancing restraining device **10** into a patient's body (an exemplary insertion step **2202**). In at least one embodiment, the restraining device **10** may be inserted through a 15 millimeter cannula under insufflation into the appropriate cavity of the patient's body. This may be achieved through use of an exemplary delivery device known in the art. At step **2202**, first swivel arm **2000** and second swivel arm **2002** of restraining device **10** are swiveled/positioned in the substantially parallel position such that the overall diameter of the restraining device **10** is sufficiently narrow for insertion into the body.

At step **2204**, restraining device **10** is advanced to a location adjacent to a targeted tissue, such as a stomach **100** (an exemplary advancement step **2204**). At step **2206** (an exemplary swivel step), first swivel arm **2000** and second swivel arm **2002** of restraining device **10** are swiveled from a substantially parallel position to a substantially perpendicular position, separating the first and second engaging components **12**, **16** from one another to a native interior space **70**. Step **2206** may be performed using any number of standard laparoscopic tools known in the art useful to pull and grasp portions of a tissue or a device. In this manner, neither the proximal ends **13**, **17** nor distal ends **14**, **18** of the first and second engaging components **12**, **16** of restraining device **10** are blocked by first swivel arm **2000** and second swivel arm **2002**, and the first and second engaging components **12**, **16** may be advanced over a targeted tissue, such as a stomach **100**.

At step **2208**, and under fluoroscopy, direct camera control or otherwise, restraining device **10** is positioned over the targeted tissue (an exemplary positioning step). In at least one embodiment, and at step **2206**, the first side **12A** of the first engaging component **12** is positioned adjacent to the desired surface of the targeted tissue and the first side **16A** of the second engaging component **16** is positioned adjacent to an opposite side of the targeted tissue. As the first and second engaging components **12**, **16** of the restraining device **10** are positioned adjacent to opposite sides of the targeted tissue, at this step **2208** the targeted tissue is positioned within the interior space **70** formed between the first and second engaging components **12**, **16**. Further, due to the configuration and composition of restraining device **10**, restraining device **10**

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can remain within the patient's body for as long as the restoration or support treatment delivered thereby is desired.

After restraining device **10** is positioned about a targeted tissue (by way of performing positioning step **2208**), method **2200** may further comprise the optional steps of securing one or more sutures to connect restraining device **10** to the targeted tissue (an exemplary suturing step **2210**), and may further comprise the step of securing cover flap **2012** to further secure restraining device about the targeted tissue (an exemplary cover flap step **2212**). In addition, and as shown in FIG. **22**, method **2200** may further comprise the step of adjusting tape **2026** to control/adjust the interior space **70** between the first engaging component **12** and the second engaging component **16** whereby at least a portion of the targeted tissue is positioned therebetween (an exemplary tape adjustment step **2214**).

In the exemplary embodiment of a restraining device **10**, as shown in FIG. **23**, restraining device **10** comprises one or more adjustment rods **2300** coupled to the first swivel arm **2000** and the second swivel arm **2002**. In the embodiment shown in FIG. **23**, for example, adjustment rods **2300** are coupled to first swivel arm **2000** and second swivel arm **2002** at or near bends **2020**, **2022**, and are further coupled to a first interconnection arm **2016** and a second interconnection arm **2018**, respectively. As shown in FIG. **23**, at in at least one embodiment, adjustment rods **2300** are coupled to first swivel arm **2000** and second swivel arm **2002** through swivel arm apertures **2302** defined therethrough, and are further coupled to first interconnection arm **2016** and second interconnection arm **2018** through interconnection arm apertures **2304** defined therethrough.

Adjustment rods **2300**, as shown in FIGS. **23** and **25**, further comprise a bar **2306** coupled to adjustment rods **2300** at or near a distal end **2308** of adjustment rod **2300**. Bar **2306**, as shown in FIG. **24**, is positioned distal to first interconnection arm **2016**/second interconnection arm **2018**, so that adjustment of a dial **2310** coupled to adjustment rod **2300** causes movement of first engaging component **12** and second engaging component **16** toward or away from one another. As shown in FIG. **24**, first interconnection arm **2016** and second interconnection arm **2018**, in various embodiments, may extend either completely or substantially along the length of first swivel arm **2000** and second swivel arm **2002** distal to first interconnection arm **2016** and second interconnection arm **2018**, so that adjustment of dial **2310** facilitates movement of first swivel arm **2000** and second swivel arm **2002**, thus facilitating movement of first engaging component **12** and second engaging component **16**.

As shown in FIGS. **23-25**, adjustment rod **2300** may further comprise a cap **2312** coupled thereto at or near a proximal end **2314** of adjustment rod **2300**. Cap **2312**, as shown in FIGS. **23-25**, may prevent dial **2310** from disengaging adjustment rod **2300** so that when restraining device **10** is positioned within a body, dial **2310** remains upon restraining device **10**. Cap **2312**, in at least one embodiment, may be bulbous/rounded at one end for comfort when restraining device **10** is positioned within a body.

As shown in FIG. **24**, dial **2310** may facilitate adjustment of device **10** by way of threads **2316** positioned along adjustment rod **2300**. In such an embodiment, rotation of dial **2310** in a first direction would cause first engaging component **12** and second engaging component **16** to move toward one another, and rotation of dial **2310** in a second/opposite direction would cause first engaging component **12** and second engaging component **16** to move away from one another. As such, restraining device **10** may be positioned within a body

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about a stomach, for example, and be adjusted using dial **2310** so that restraining device **10** is optimally positioned about the stomach.

As shown in the exemplary embodiments in FIGS. **25** and **26**, dial **2310** may define a dial aperture **2318**, whereby indicia **2320** upon adjustment rod **2300** may be viewed therethrough. As shown in FIG. **26**, for example, as dial **2310** is rotated, one or more indicia **2320** may be visible through dial aperture **2318**, so that a user of such an embodiment of a restraining device **10** of the present disclosure may identify a level of adjustment. Indicia **2320** may be one or more numbers, letters, lines and/or other indicia **2320** useful to identify a level of adjustment of restraining device **10**.

As shown in FIG. **27**, and in at least one embodiment of a restraining device **10** of the present disclosure, restraining device **10** further comprises at least one flange **2700** coupled to first swivel arm **2000** and/or second swivel arm **2002**. Flange **2700** defines an aperture therethrough so that adjustment rod **2300** may fit therethrough, providing physical support for dial **2310** as dial **2310** engages said flange **2700**.

An exemplary embodiment of a restraining device **10** of the present disclosure having adjustment rods **2300** is shown in FIG. **28**. As shown in FIG. **28**, first engaging component **12** and/or second engaging component **16** may define one or more facets **2800** along at least part of the length of said components **12**, **16**, whereby facets **2800** provide a generally arcuate profile as shown therein. Facets **2800** may further improve the overall comfort of restraining device **10** when it is positioned with a patient's body.

Steps of a method for delivering and positioning an embodiment of a restraining device **10**, such as the embodiment shown in FIG. **23**, are shown in FIG. **29**. As shown in FIG. **29**, exemplary method **2900** comprises the step of laparoscopically inserting/advancing restraining device **10** into a patient's body (an exemplary insertion step **2202**) as previously discussed herein. At step **2202**, first swivel arm **2000** and second swivel arm **2002** of restraining device **10** are swiveled/positioned in the substantially parallel position such that the overall diameter of the restraining device **10** is sufficiently narrow for insertion into the body.

At step **2204**, restraining device **10** is advanced to a location adjacent to a targeted tissue, such as a stomach **100** (an exemplary advancement step **2204**). At step **2206** (an exemplary swivel step), first swivel arm **2000** and second swivel arm **2002** of restraining device **10** are swiveled from a substantially parallel position to a substantially perpendicular position, separating the first and second engaging components **12**, **16** from one another to a native interior space **70** as previously discussed herein.

At step **2208**, and under fluoroscopy, direct camera control or otherwise, restraining device **10** is positioned over the targeted tissue (an exemplary positioning step) as previously referenced herein. After restraining device **10** is positioned about a targeted tissue (by way of performing positioning step **2208**), method **2900** may further comprise the optional steps of securing one or more sutures to connect restraining device **10** to the targeted tissue (an exemplary suturing step **2210**), and regarding an embodiment of restraining device **10** of the present disclosure comprising a cover flap **2012**, method **2900** may further comprise the step of securing cover flap **2012** to further secure restraining device about the targeted tissue (an exemplary cover flap step **2212**). In addition, and as shown in FIG. **29**, method **2900** may further comprise the optional step of rotating dial **2310** to control/adjust the interior space **70** between the first engaging component **12** and the second engaging component **16** whereby at least a portion

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of the targeted tissue is positioned therebetween (an exemplary dial adjustment step 2902).

The various embodiments of restraining devices 10 described herein provide numerous benefits over the devices and systems of the prior art. An exemplary restraining device 10 may be inserted laparoscopically and/or endoscopically, is minimally invasive, completely reversible and available for chronic placement without the risk of complications. Furthermore, use of a restraining device 10 to treat and/or support a targeted tissue or organ produces a reduced amount of negative side effects than the procedures of the prior art for similar indications. While embodiments of restraining devices 10 are presented with respect to specific anatomy and treatment examples, the various restraining devices 10 and methods 300 may be expanded for use in treating any organ, limb or body structure that would benefit from reshaping, restoring, or added support provided through a reversible, easy to use and easy to implement technique for chronic placement.

Prior art focuses on creating a restrictive line through “clamps” to mimic the gastric sleeve. The disclosure of the present application provides various loosely-fitting restraints (to prevent migration) and to affect motility of the stomach. The contractility or motility curve of the gastric tissue is substantially attenuated by an external restraint (as discussed below), thus affecting the movement of bolus through the stomach. Hence, the food volume will remain in the pouch longer and lead to earlier distension which affects neuroactivity and mechanosensory elements of satiety. Hence, the principle of the restraining devices 10 of the present disclosure is based on flow (motility) unlike previous art that focuses on restrictive which risks migration and erosion.

As described herein, placement of exemplary restraining devices 10 through the lower curvature can be easily implemented as opposed to procedures which approach the greater curvature where the liver and diaphragm are in the proximity. Furthermore, various clips, clamps, or perforated plates known in the art which transverse the stomach (as opposed to sleeve) can lead to substantial remodeling of the fundus which would lead to weight regain after a period of time. As such, the various connectors of the disclosure of the present application operate to prevent overall stretch of stomach and hence prevent remodeling. The stretch that leads to mechanosensory satiety in the restraining devices of the present disclosure occurs locally between the restraining springs or bars while still restraining the overall pouch. No such global containment of the gastric tissue is known in the prior art. Finally, various clips, clamps, or perforated plates known in the art would exert significant local stresses on the tissue which can lead to erosion or migration as the substantially restricted stomach would attempt to distend. Conversely, the first engaging component 12 and the second engaging component 16 of the restraining devices 10 of the present disclosure distribute the stress more uniformly over the plate and only restrain a relatively small pouch rather than a more substantial portion of the stomach transversely.

While various embodiments of implantable restraining devices, systems, and methods for using the same have been described in considerable detail herein, the embodiments are merely offered by way of non-limiting examples of the disclosure described herein. It will therefore be understood that various changes and modifications may be made, and equivalents may be substituted for elements thereof, without departing from the scope of the disclosure. Indeed, this disclosure is not intended to be exhaustive or to limit the scope of the disclosure.

Further, in describing representative embodiments, the disclosure may have presented a method and/or process as a

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particular sequence of steps. However, to the extent that the method or process does not rely on the particular order of steps set forth herein, the method or process should not be limited to the particular sequence of steps described. Other sequences of steps may be possible. Therefore, the particular order of the steps disclosed herein should not be construed as limitations of the present disclosure. In addition, disclosure directed to a method and/or process should not be limited to the performance of their steps in the order written. Such sequences may be varied and still remain within the scope of the present disclosure.

The invention claimed is:

1. An implantable restraining device, comprising:

a first engaging component and a second engaging component defining a longitudinal axis, each component configured for laparoscopic insertion into a body cavity, wherein the first engaging component and the second engaging component each comprise a proximal end, a distal end, and a body extending therebetween, wherein the body of the first engaging component is configured to conform to a first targeted tissue surface, and the body of the second engaging component is configured to conform to a second targeted tissue surface;

a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point; and

a cover flap, the cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging component when initially coupled to the second engaging component; and

wherein the first swivel arm and the second swivel arm are configured to move from a first position that is substantially coplanar with the first engaging component and the second engaging component to a second position that is substantially perpendicular with the first engaging component and the second engaging component so to open the device, said movement in a plane perpendicular to a plane defined by an interior space between the first engaging component and the second engaging component;

and wherein when the first swivel arm and the second swivel arm are not positioned within the interior space between the first engaging component and the second engaging component when in the second position when the device is open.

2. The device of claim 1, wherein the first engaging component and the second engaging component are configured to engage a targeted tissue therebetween when the first swivel arm and the second swivel arm are in a configuration relatively perpendicular to the first engaging component and the second engaging component.

3. The device of claim 1, wherein the first engaging component and the second engaging component each define one or more studs sized and shaped to permit the first swivel arm and the second swivel arm to be coupled thereto.

4. The device of claim 1, wherein the first swivel arm and the second swivel arm further define one or more curvatures, whereby the first bend, the second bend, and the one or more curvatures define a native U-shaped configuration.

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5. The device of claim 1, further comprising:
 a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend; and
 a tape positioned around at least part of the device so that the tape engages the first interconnection arm and the second interconnection arm.
6. The device of claim 5, wherein the tape is capable of decreasing an interior space defined between the first engaging component and the second engaging component when the tape applies a force to the first interconnection arm and the second interconnection arm.
7. The device of claim 5, wherein the tape further comprises one or more detectable portions thereon.
8. The device of claim 7, wherein the one or more detectable portions are radiopaque.
9. The device of claim 1, wherein at least one of the first engaging component and the second engaging component define one or more suture apertures therethrough.
10. The device of claim 1, wherein each of the first engaging component and the second engaging component define one or more facets along at least part of a length of said engaging components, said facets providing a generally arcuate profile of said engaging components.
11. The device of claim 1, wherein each of the first engaging component and the second engaging component comprise a configuration chosen from a straight bar configuration, a curved configuration, or a circular configuration.
12. The device of claim 1, wherein the first engaging component and the second engaging component are flexible or semi-flexible.
13. The device of claim 1, wherein the first engaging component, the second engaging component, the first swivel arm, and the second swivel arm each comprise a material suitable to resist corrosion selected from the group consisting of polyurethane, polyethylene, polytetrafluoroethylene, nitinol, silastic, titanium, and stainless steel.
14. The device of claim 1, wherein the first engaging component, the second engaging component, the first swivel arm, and the second swivel arm comprise a resorbable material selected from the group consisting of polyglycolide (PGA), polylactide (PLA), L-lactide (LPLA), poly(DL-lactide) (DL-PLA), poly(ϵ -caprolactone) (PCL), poly(dioxanone) (PDO), polyglycolide-trimethylene carbonate (PGA-TMC), or poly(D,L-lactide-co-glycolide) (DLPLG).
15. An implantable restraining device, comprising:
 a first engaging component and a second engaging component, each component configured for laparoscopic insertion into a body cavity;
 a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point;
 a first interconnection arm and a second interconnection arm, the first interconnection arm connected to the first swivel arm relative to the first bend, and the second interconnection arm connected to the second swivel arm relative to the second bend;
 a tape positioned around at least part of the device so that the tape engages the first interconnection arm and the second interconnection arm; and

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- a cover flap, the cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging component when initially coupled to the second engaging component;
 wherein the first swivel arm and the second swivel arm are capable of moving between a first position that is substantially parallel with the first engaging component and the second engaging component and a second position that is substantially perpendicular with the first engaging component and the second engaging component.
16. The device of claim 15, wherein the first engaging component and the second engaging component are configured to engage a targeted tissue therebetween when the first swivel arm and the second swivel arm are in a configuration relatively perpendicular to the first engaging component and the second engaging component.
17. The device of claim 15, wherein the tape is capable of decreasing an interior space defined between the first engaging component and the second engaging component when the tape applies a force to the first interconnection arm and the second interconnection arm.
18. The device of claim 15, wherein the first engaging component and the second engaging component are configured to engage a targeted tissue therebetween when the first swivel arm and the second swivel arm are in a configuration relatively perpendicular to the first engaging component and the second engaging component.
19. The device of claim 15, wherein the first swivel arm and the second swivel arm further define one or more curvatures, whereby the first bend, the second bend, and the one or more curvatures define a native U-shaped configuration.
20. The device of claim 15, wherein at least one of the first engaging component and the second engaging component define one or more suture apertures therethrough.
21. The device of claim 15, wherein each of the first engaging component and the second engaging component comprise a configuration chosen from a straight bar configuration, a curved configuration, or a circular configuration.
22. The device of claim 15, wherein the first engaging component, the second engaging component, the first swivel arm, and the second swivel arm each comprise a material suitable to resist corrosion selected from the group consisting of polyurethane, polyethylene, polytetrafluoroethylene, nitinol, silastic, titanium, and stainless steel.
23. An implantable restraining device, comprising:
 a first engaging component and a second engaging component defining a longitudinal axis, each component configured for laparoscopic insertion into a body cavity;
 a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point; and
 a cover flap, the cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging component when initially coupled to the second engaging component;

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wherein the first engaging component and the second engaging component each define one or more studs sized and shaped to permit the first swivel arm and the second swivel arm to be coupled thereto;

wherein the first swivel arm and the second swivel arm are configured to move from a first position that is substantially coplanar with the first engaging component and the second engaging component to a second position that is substantially perpendicular with the first engaging component and the second engaging component so to open the device, said movement in a plane perpendicular to a plane defined by an interior space between the first engaging component and the second engaging component;

and wherein when the first swivel arm and the second swivel arm are not positioned within the interior space between the first engaging component and the second engaging component when in the second position when the device is open.

24. The device of claim **23**, wherein the first engaging component and the second engaging component are configured to engage a targeted tissue therebetween when the first swivel arm and the second swivel arm are in a configuration relatively perpendicular to the first engaging component and the second engaging component.

25. The device of claim **23**, wherein the first swivel arm and the second swivel arm further define one or more curvatures, whereby the first bend, the second bend, and the one or more curvatures define a native U-shaped configuration.

26. The device of claim **23**, wherein at least one of the first engaging component and the second engaging component define one or more suture apertures therethrough.

27. The device of claim **23**, wherein the first engaging component and the second engaging component are flexible or semi-flexible.

28. A method for restoring a targeted tissue, the method comprising the steps of:

inserting an implantable restraining device into a body cavity of a mammalian body, the implantable restraining device comprising:

a first engaging component and a second engaging component defining a longitudinal axis, each component configured for laparoscopic insertion into a body cavity, wherein the first engaging component and the second engaging component each comprise a proximal end, a distal end, and a body extending therebetween, wherein the body of the first engaging component is configured to conform to a first targeted tissue surface, and the body of the second engaging component is configured to conform to a second targeted tissue surface,

a first swivel arm defining a first bend and a second swivel arm defining a second bend, the first swivel arm coupled to the first engaging component at a first pivot point and coupled to the second engaging component at a second pivot point, and the second swivel arm coupled to the

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first engaging component at a third pivot point and coupled to the second engaging component at a fourth pivot point, and

a cover flap coupled to either the first engaging component or the second engaging component, the cover flap capable of either further coupling to the second engaging component when initially coupled to the first engaging component or further coupling to the first engaging component when initially coupled to the second engaging component,

wherein the first swivel arm and the second swivel arm are configured to move from a first position that is substantially coplanar with the first engaging component and the second engaging component to a second position that is substantially perpendicular with the first engaging component and the second engaging component so to open the device, said movement in a plane perpendicular to a plane defined by an interior space between the first engaging component and the second engaging component;

wherein when the first swivel arm and the second swivel arm are not positioned within the interior space between the first engaging component and the second engaging component when in the second position when the device is open;

advancing the implantable restraining device to a location within the mammalian body adjacent to a targeted tissue;

swiveling the first swivel arm and the second swivel arm so that the first swivel arm and the second swivel arm are substantially perpendicular to the first engaging component and the second engaging component; and

positioning the first engaging component and the second engaging component over the targeted tissue such that at least a portion of the targeted tissue is positioned therebetween, wherein when the targeted tissue expands in a direction between the first engaging component and the second engaging component, the targeted tissue exerts a force upon the first engaging component and the second engaging component.

29. The method of claim **28**, further comprising the step of: securing one or more sutures to connect the first engaging component and/or the second engaging component to the targeted tissue.

30. The method of claim **28**, further comprising the step of: securing the cover flap so that the cover flap is secured to the first engaging component and the second engaging component.

31. The method of claim **28**, wherein the targeted tissue is a stomach, and wherein expansion of the stomach, with said device positioned thereon, functionally divides the stomach into a first stomach portion and a second stomach portion.

32. The method of claim **28**, further comprising the step of: securing the cover flap over a fundus of the stomach so that the cover flap is secured to the first engaging component and the second engaging component.

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